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- (71) Applicant THE BOARD OF REGENTS OF THE UNIVERSITY OF OKLAHOMA Norman, Oklahoma 73109 (US)
- (72) Inventors.
 Weigel, Paul H.
 Edmond, OK 73063 (US)

- DeAngelis, Paul
- Edmond, OK 73034 (US)
- Kumari, Kshama Edmond, OK 73013 (US)
- (74) Representative: Ebner von Eschenbach, Jennifer et al Ladas & Parry, Dachauerstrasse 37 80335 München (DE)

Remarks:

This application was filed on 09 - 12 - 2004 as a divisional application to the application mentioned under INID code 62.

- (54) Hyaluronan synthase gene and uses thereof
- (57) An isolated enzymatically-active hyaluronan synthase (HAS) is provided which is encoded by a nucleic acid sequence that has at least 80% identity with SEQ ID NO: 8. In particular, the HAS is a chlorella virus polypeptide. A nucleic acid segment encoding the HAS

is also provided, which nucleic acid segment is used in the preparation of recombinant cells which produce hyaluronate synthase and its hyaluronic acid product. Hyalurante is also known as hyaluronicacid or hyaluronan.

Description

BACKGROUND OF THE INVENTION

5 1 Field of the Invention

[0001] The present invention relates to a nucleic acid segment having a coding region segment encoding enzymaticulty active Streptococcus equisimilis hyaluronate synthase (seHAS), and to the use of this nucleic acid segment in the preparation of recombinant cells which produce hyaluronate synthase and its hyaluronic acid product. Hyaluronate is also known as hyaluronic acid or hyaluronan.

2. Brief Description of the Related Art.

[0002] The incidence of streptococcal infections is a major health and economic problem worldwide, particularly in developing countries. One reason for this is due to the ability of Streptococcal bacteria to grow undelected by the body's phagocytic cells, i.e., macrophages and polymorphonuclear cells (PMNs). These cells are responsible for recognizing and engulfing foreign microorganisms. One effective way the bacteria evade surveillance is by coating thereselves with polysaccharide capacies, such as a hypluronia cell (HA) capacie. The structure of HA is identical inhorth prokaryotes and eukaryotes. Since HA is generally nonimmunogenic, the encapsulated bacteria do not elicit an immuneresponse and are, therefore, not targreted for destruction. Moreover, the capacite everts an antihipsacycilo effect on PMNs in vitro and prevents ettachment of Streptococcus to macrophages. Precisely because of this, in Group A and Group C Streptococci, the HA capacites are major virulence factors in natural and experimental infections. Group A Streptococcus are responsible for numerous human diseases including pharpropilis, impetigo, deep issue infections, therefore, the proping the properties of the proping the propi

[003] Structurally, HA is a high molecular weight linear polysaccharide of repeating disaccharide units consisting of N-acetylgilucosamine (GloHa). The number of repeating disaccharides in an HA molecule can exceed 30.000, a M>10.7 HA is the only glycosaminogylcan synthesized by both mammalian and bacterial cells particularly Groups A and C Streptococci and Type A Pasturella multicade. These strains make HA which is secretal into the medium as well as HA is of broad Interest medicinally since the production of the HA capsule is a very efficient and clever way that Streptococci use to evade surveillance by the immune system.

[004] HA is synthesized by mammalian and bacterial cells by the enzyme hyaluronate synthase which has been localized to the plasma membrane. It is believed that the synthesis of HA in these organisms is a multi-step process, initiation involves binding of an initial precursor, UDP-GlorNAc or UDP-GlorA. This is followed by elongation which involves alternate addition of the two sugars to the, growing oligosaccharide chain. The growing polymer is extruded across the plasma membrane region of the cell and into the extracellular space. Although the HA biosynthetic system was one of the first membrane heteropolysaccharide synthetic pathways studied, the mechanism of HA synthesis is still not well understood. This may be because in vitro systems developed to date are inadequate in that de novo biosynthesis of HA has not been accomplished.

[0005] The direction of HA polymer growth is still a matter of disagreement among those of ordinary Skill in the art.

Addition of the monosaccharides could be to the reducing or nonreducing end of the growing HA chain. Furthermore, questions remain concerning (i) whether chairs are initiated using a primer, and (iii) the mechanism by which the mature polymer is ectuded through the plasma membrane of the Streptococcus. Understanding the mechanism of HA biosynthesis may allow development of alternative strategies to control Streptococcus and Peutrel'is infections by interfering in the process [0006] HA has been identified in virtually every tissue in vertebrates and has achieved widespread use in various clinical applications, most notably and appropriately as an intra-articular matrix supplement and in eye surgery. The scientific literature has also shown a transition from the original perception that HA is primarily a passive structural component in the matrix of a few connective tissues and in the capsule of cortain strains of bactera to a recognition that this ubiquitious macromiecules indynamically involved in many biological processes. From modulating cell migration and differentiation during embryogenesis to regulation of extracellular matrix organization and metabolism to important roles in the complex processes of metabasis, wound healing, and inflammation. Further, it is becoming clear that HA is highly metabolically active and that cells focus much attention on the processes of its synthesis and catabolism. For example, the half-life of HA in tissues ranges from 1 to 3 weeks in cartilage to <1 day in repidermis.

[0007] It is now clear that a single protein utilizes both sugar substrates to synthesize HA. The abbreviation HAS, for the HA synthase, has gained videspread support for designating this class of enzymes. Markovit et al. successfully characterized the HAS activity from Streptococcus viocenes and discovered the enzymes for membrane localization.

and its requirements for sugar nucleotide precursors and Mi²² Prehm found that elongating HA, made by B6 cells, was digested by hyaluronidase added to the medium and proposed that HAS resides at the plasma membrane. Philipson and Schwartz also showed that HAS activity cofractionated with plasma membrane markers in mouse oligoden-droolioms cells.

[0008]. HAS assembles high M, HA that is simultaneously extruded through the membrane into the extracellular space (or to make the cell capsule in the case of bacteria) as glycosaminoglycan synthesis proceeds. This mode of biosynthesis is unique among macromolecules since nucleic acids, proteins, and lipids are synthesized in the nucleus, endoplasmic reticulum/Golgi, cytoplasm, or mitochondria. The extrusion of the growing chain into the extracellular space also allows for unconstrained polymer growth, thereby activering the exceptionally large size of HA, whereas confinement of synthesis within a Golgi or post-Golgi compartment could limit the overall amount or length of the polymers formed. High concentrations of HA within a confined lumen could also create a high viscosity environment that might be deleterious for other organically functions.

[0009] Several studies attempted to solubilize, identify, and purify HAS from strains of Streptococci that make a capsular coat of HA as well as from eukaryotic cells. Although the streptococcal and murino oligodendroglioms enzymes were successfully detergent-solubilized and studied, efforts to purify an active HAS for further study or molecular cloning remained unsuccessful for decades. Prehm and Mausoff used periodate-oxidized UDP-Glock or UDP-GlockA to effinite pale al protein of ~52 KD as it explococcal membranes that co-purified with HAS. This led to a report claiming that the Group C streptococcal HAS had been cloned, which was unfortunately erroneous. This study failed to demonstrate expression of an active synthase and may have actually cloned a peptide transporter. Tiscott and van de Rijn used digitonin to solubilize HAS from streptococcal membranes in an active form. Van de Rijn and Drake selectively radiolabeled three streptococcal membrane proteins of 42, 33, and 27 KDa with 5-actio-UDP-Glock and suggested that the 33-kDa protein was HAS. As shown alse, however, HAS actually turned out to be the 42-kDa protein.

[0010] Despite these efforts, progress in understanding the regulation and mechanisms of HA synthesis was essentially stalled, since there were no molecular probes for HAS mRNA or HAS protein. A major breakthrough occurred in 1993 when DeAngelis of all reported the molecular cloning and characterization of the Group A streptococcal gene encoding the protein HasA. This gene was known to be in part of an operon required for bacterial HA synthesis, although the function of this protein, which is now designated as spiHAS (the S. progenes HAS), was unknown. spiHAS was subsequently proven to be responsible for HA elongation and was the first glycosaminoglycan synthase dentified and cloned and then successfully expressed. The S. progenes HA synthesis operon encodes two other proteins. HasB is a UDP-glucose dehydrogenase, which is required to convert UDP-glucose to UDP-GlucA- one of the substrates for HA synthesis. HasC is a UDP-glucose pyrophosphorylase, which is required to convert glucose 1-phosphate and UTP to UDP-glucose Co-transfection of both hasA and hasB genes into either acapsular Streptococcus strains or Enterous faecalis conferred them with the ability to synthesize HA and form a capsule. This provided the first strong evidence that HasA is an HA synthesis.

[0011] The elusive HA synthase gene was finally cloned by a transposon mutagenesis approach, in which an acapsular mutant Group A strain was created containing a transposon interruption of the HA synthesis operon. Known
sequences of the transposon allowed the region of the junction with streptococcal DNA to be identified and then cloned
from wild-type cells. The encoded sph4S was 5-10% identical to a family of yeast-chitin synthases and 30% identical
to the Xenopus leavis protein DG42 (developmentally expressed during gastrulation), whose function was unknown
at the time. DeAngelis and Vieigel expressed the active recombinant sph4S in Escherichia coll and showed that this
single purified gene product synthesizes high Mi, HA when inclushed in vitro with UDP-GIGA And DIP-GIGNAC. Thereby
showing that both glycosyltransferase activities required for HA synthesis are catalyzed by the same protein, as first
proposed in 1998. This set the stage for the almost simultaneous identification of eukaryotic HAS CINAs in 1996 by
four laboratories revealing that HAS is a multigene family encoding distinct isozymes. Two genes (HAS? and HAS2)
were quickly discovered in memmals (20-34), and a third gene HAS3 was stard efiscovered. A second sterptocoal
seHAS or Streptococcus equisimilis hyaluronate synthase, has now been found and is the invention being claimed
and disclosed herein.

[0012] As indicated, we have also identified the authentic HAS gene from Group C Streptococcus equisimitis (se-HAS), the seHAS protein has a high level of identity (approximately 70 percent) to the spHAS gene.

This identity, however, is interesting because the seHAS gene does not cross-hybridize to the spHAS gene.

[0013] Membranes prepared from E. cotl expressing recombinant setHAS synthesize HA when both substrates are provided. The results confirm that the earlier report of Lansing et al. claiming to have cloned the Group C HAS we wrong. Unfortunately, several studies have employed antibody to this uncharacterized 52-kDa streptococcal protein to investigate what was believed to be eukaryotic HAS.

[0014] Itano and Kimata used expression cloning in a mutant mouse mammary carcinoma cell line, unable to synthesize HA, to clone the first putative mammalian HAS cDNA (mmHAST). Subclones defective in HA synthesis fell into three separate classes that were complementary for HA synthesis in somatic cell fusion experiments, suggesting that at least three proteins are required. Two of these classes maintained some HA synthetic activity, whereas one

showed none. The latter cell line was used in transient transfection experiments with cDNA prepared from the parental cells to identify a single protein that restored HA synthetic activity. Sequence analyses revealed a deduced primary structure for a protein of -65 kDa with a predicted membrane topology similar to that of spirAS. mmHAS1 is 30% identical to 5pHAS and 55% identical to UG42. The same month this report appeared, bries other groups submitted papers describing cDNAs encoding what was initially thought to be the same mouse and human enzyme. However, through an extraordinary circumstance, each of the four laboratories had discovered a separate HAS isozyme in both species.

[0015] Using a similar functional cloning approach to that of Itano and Kimata, Shyjan et al. identified the human homolog of HAS 1. A mesenteric lymph node cDNA library was used to transfect mutine mucosal T lymphocytes that were then screened for their ability to adhere in a rosette assay. Adhesion of one transfectant was inhibited by antisera to CD44, a known cell surface HA-kinding protein, and was abrogated directly by pretreatment with hyaturonidase. Thus, rosetting by this transfectant required synthesis of HA Cloning and sequencing of the responsible CDNA indicated hsHAS1, Itano and Kimata also reported a human HAS1 cDNA solated from a fetal brain library. The hsHAS1 cDNAs reported by the two groups, however, differ in length; they encode a 578 or a 543 amino acid protein. HAS activity has only been demonstrated for the longer form.

[0016] Based on the molecular identification of spHAS as an authentic HA synthase and regions of near identity among DG42 spHAS, and NodC (a β-GloNAc transferase nodulation factor in Rhizobium). Spicer et al. used a degenerate RT-PCR approach to clone a mouse embryo cDNA encoding a second distinct enzyme, which is designated mmHAS2. Transfection of mmHAS2 aDNA into COS cells directed de novo production of an HA cell cot actie etail military approach. Watanabe and Yamaguchi screened a human fetal brain cDNA library to identify hat-RS2. Fulce et al. independently used a similar strategy to identify mmHAS2 in RNA isolated from ovarian cumulus cells-cocyte complexes were isolated from mice immediately after initiating an ovulatory cycle, before HA synthesia to at later times when HA synthesia is just beginning (3 h) or already apparent (4 h). RT-PCR showed that HAS2 a mRNA was absent initially but expressed at high levels 2-4 h later suggesting that transcription of HAS2 regulates HA synthesis in this process. Both hat-MS2 are 552 amino acids in length and are 98% identical, mmHAS1 is 583 amino acids long a 95% identical to hat-RS1, which is 578 amino acids in length and are 98% identical, mmHAS1 is 583 amino acids long a 95% identical which is 578 amino acids in length and are 98% identical.

[0017]. Most recently Spicer et al. used a PCR approach to identify a third HAS gene in mammals. The mmHASS protein is 554 amino acids long and 71, 56, and 28% identical, respectively, to mmHAS1, mmHAS2, DG42, and spHAS Spicer et al. have also localized the three human and mouse genes to three different chromosomes (HAS1 to hsChr 19/mmChr 17; HAS2 to hsChr 8/mmChr 15; HAS3 to hsChr 8/mmChr 16; Lacilization of the three HAS genes on different chromosomes and the appearance of HA throughout the vertebrate class suggest that this gene family is ancient and that isozymes appeared by duplication early in the evolution of vertebrates. The high identity (~20%) between the bacterial and eukaryotic HAS also suggests that the von had a common ancestral gene. Perhaps primitive bacteria usurped the HAS gene from an early vertebrate when the deciral are products became larger and more complex. Alternatively, the bacteria could have obtained a larger vertebrate HAS gene and deleted regulatory sequences nonessential for enzyme activity.

[0018] The discovery of X. Iaevis DG42 by Dawid and co-workers played a significant role in these recent developments, even though this protein was not known to be an HA synthase. Nonetheless, that DG42 and spHAS were 30% identical was critical for designing oligonucleotides that allowed identification of mammalian HAS2. Inonically, definitive evidence that DG42 is a bone fide HA synthase was reported only after the discoveries of the Mammalian isozymes, when DeAngleis and Achytuthan expressed the recombinant protein in yeast (an organism that cannot synthesize) when DeAngleis and Achytuthan expressed the recombinant protein in yeast (an organism that cannot synthesize also showed that it synthesizes HA when isolated membranes are provided with the two substrates. Meyer and Kreil also showed that Iysates from cells transfected with cDNA for DG42 synthesize elevated levels of HA. Now that its function is known, DG42 can, therefore, be designated X1HAS.

[0019] There are common predicted structural features shared by all the HAS proteins, including a large central domain and clusters of 2-3 transmembrane or membrane-associated domains at both the amino and carboysl ends of the protein. The central domain, which comprises up to ~88% of the predicted intracellular HAS protein sequences, probably contains the catalytic regions of the enzyme. This predicted central domain is 264 amino acids long in spiHAS (63% of the total protein) and 07-328 residues long in the eukaryotic HAS members (64-65% of the total protein). The exact number and orientation of membrane domains and the topological organization of extracellular and intracellular looss have not vert been experimentally determined for any HAS

[0020] spHAS is a HAS family member that has been purified and partially characterized. Initial studies using spHAS/ sikaline phosphatase fusion proteins indicate that the N terminus, C terminus, and the large central domain of spHAS are, in fact, inside the cell: spHAS has 6 cysteines, whereas HAS1, HAS2, and HAS3 have 13, 14 and 14 Cys residues, respectively. Two of the 6 Cys residues in spHAS are conserved and identical in HAS1 and HAS2. Only one conserved Cys residue is found at the same position (Cys-25 in spHAS) in all the HAS family members. The may be an essential

Cys whose modification by sufflydryl poisons partially inhibits enzyme activity. The possible presence of disulfide bonds or the identification of critical Cys residues needed for any of the multiple HAS functions noted below has not yet been elucidated for any members of the HAS family.

[0021] In addition to the proposed unique mode of synthesis at the plasma membrane, the HAS enzyme family is highly unusual in the large number of functions required for the overall polymerization of HA. At least six discrete activities are present within the HAS enzyme; binding sitse for each of the two different sugar nucleotide precursors (UDP-GloNAc and UDP-GloA), two different glycosyltransferase activities, one or more binding sitse that anchor the growing HA polymer to the enzyme (perhaps related to a B-X₇-B motif), and a ratiohet-like transfer reaction that moves the growing polymer one sugar at a time. This later activity is likely coincident with the stepwise advance of the polymer through the membrane. All of these functions, and perhaps others as yet unknown, are present in a relatively small protein rainign in size from 419 (spHAS) to S68 (trHAS) amino acids.

[0022] Although all the available evidence supports the conclusion that only the spHAS protein is required for HA biosynthesis in bacters or in vitro, it is possible that the larger euklaryotic HAS lamily embers are part of multicomponent complexes. Since the euklaryotic HAS proteins are -40% larger than spHAS, their additional protein dominacould be involved in more elaborate functions such as intracellular trafficking and localization, regulation of enzyme activity, and mediating interactions with other cellular components.

[0023] The unexpected finding that there are multiple vertebrate HAS genes encoding different synthases strongly supports the emerging consensus that HA is an important regulator of cell behavior and not simply a structural component in fessues. Thus, in less than six months, the field moved from one known, cloned HAS (spHAS) to recognition of a multigene family that promises rapid, numerous, and exciting future advances in our understanding of the synthesis and biology of HA.

[0024] For example, disclosed hereinsafter are the sequences of the two HAS genes: from Pasturells multiplications and (2) Paramecium bursaria chlorelle virus (PBCV-1). The presence of hyaturonan synthase in these two systems and the purification and use of the hyaturonan synthase from these two different systems indicates an ability to purify and isolate nucleic acid sequences encoding enzymatically active hyaturonan synthase in many different prokaryotic and viral sources.

[0025] Group C Streptococcus equisimilis strain D181 synthesizes and secretes hyaluronic acid (HA). Investigators have used this strain and Group A Streptococcus grogone strains, such as S43 and A111, to study the biosynthesis of HA and to characterize the HA-synthesizing activity in terms of its divalent cation requirement, precursor (UDP-Gic-NAc and UDP-GicA) utilization, and obtimum bH.

[9028] Traditionally, IAA has been prepared commercially by isolation from either rooster combs or extracellular media from Streptococcal cultures. One method which has been developed for preparing IAA is through the use of cultures of HA-producing Streptococcal bacteria. U.S. Patent No. 4,517,295 describes such a procedure wherein HA-producing Streptococcal are fermented under anaerobic conditions in a CO₂-enriched growth medium. Under these conditions, HA is produced and can be extracted from the brottl. It is generally left that isolation of HA from rooster combs is laborious and difficult, since one starts with HA in a less pure state. The advantage of isolation from rooster combs is that the HA produced is of higher molecular weight. However, preparation of HA by bacterial fermentation is easier, since the HA is of higher purity to start with. Usually, however, the molecular weight of HA produced in this way is smaller than that from rooster combs. Therefore, a technique that would allow the production of high molecular weight HA by bacterial fermentation would be an improvement over existing procedures.

[0027] High molecular weight HA has a wide variety of useful applications – ranging from cosmetics to eye surgery. Due to its potential for high viscosity and its high biocompatibility. HA finds particular application in eye surgery as a replacement for vitreous fitud. HA has also been used to treat racehorese for traumatic arthrifts by intra-articular injections of HA. In shaving cream as a lubricant, and in a variety of cosmetic products due to its physicochemical properties of high viscosity and its ability to retain moisture for long periods of time. In fact, in August of 1997 the U.S. Food and Drug Agency approved the use of high molecular weight HA in the treatment of severe arthrifts through the injection of such high molecular weight HA directly into the affected joints. In general, the higher molecular weight HA that is employed the better. This is because HA solution viscosity increases with the average molecular weight of the individual HA polymer molecules in the solution. Unfortunately, very high molecular weight HA, such as that ranging up to 10? has been difficult to obtain by currently available isolation procedures.

[0023] To address these or other difficulties, there is a need for new methods and constructs that can be used to produce HA having one or more improved properties such as greater purity or ease of preparation. In particular, there is a need to develop methodology for the production of larger amounts of relatively high molecular weight and relatively pure HA than is currently commercially available. There is yet another need to be able to develop methodology for the production of HA having a modified size distribution (HA_{Nova}) as well as HA having a modified size distribution (HA_{Nova}) as well as HA having a modified size distribution (HA_{Nova}) as well as HA having a modified size distribution (HA_{Nova}).

[0029] The present invention addresses one or more shortcomings in the art. Using recombinant DNA technology, a purified nucleic acid segment having a coding region encoding enzymatically active setHAS is disclosed and claimed in conjunction, with methods to produce an enzymatically active HA synthase, as well as methods for using the nucleic

acid segment in the preparation of recombinant cells which produce HAS and its hyaturonic acid product

[0030] Thus, it is an object of the present invention to provide a purified nucleic acid segment having a coding region encoding enzymatically active HAS.

[0031] It is a further object of the present invention to provide a recombinant vector which includes a purified nucleic acid segment having a coding region encoding enzymatically active HAS.

[0032] It is still a further object of the present invention to provide a recombinant host cell transformed with a recombinant vector which includes a purified nucleic acid segment having a coding region encoding enzymatically active the combinant vector which includes a purified nucleic acid segment having a coding region encoding enzymatically active the [18].

[0033] It is yet another object of the present invention to provide a method for detecting a bacterial cell that expresses

[0034] It is another object of the present invention to provide a method for producing high and/or low molecular weight hyaluronic acid from a hyaluronate synthase gene, such as seHAS, as well as methods for producing HA having a modified size distribution and/or a modified size this tubins and/or a modified size.

[0035] These and other objects of the present invention will become apparent in light of the attached specification, claims, and drawings

BRIEF SUMMARY OF THE INVENTION

[9038] The present invention involves the application of recombinant DNA technology to solving one or more problems in the art of hyaliuronic acid (HA) preparation. These problems are addressed through the isolation and use of a nucleic acid segment having a coding region encoding the enzymetically active Streptococcus equientilis (sel+AS) hyaliuronate synthase gene, a gene responsible for HA chain biosynthesis. The sel+AS gene was cloned from DNA of an appropriate microbial source and engineered into useful recombinant constructs for the preparation of HA and for the preparation of large quantities of the HAS enzyme itself.

[0037] The present invention encompasses a novel gene, seHAS. The expression of this gene correlates with virulence of Streptococcal Group A and Group C strains, by providing a means of escaping phagocytosis and immunes unveillance. The terms "hyalutronia cuid synthases," hyalutroniate synthase, "shutronian synthases" and "HA synthianes, are used interchangeably to describe an enzyme that polymerizes a glycosaminoglycan polysaccharide chain composed of alternating glucuronic acid and N-acetylglucosamine sugars, β 1,3 and β 1,4 linked. The term "seHAS" describes the HAS enzyme defined from Streptococcus equisiminis.

[0038] The present invention concerns the isolation and characterization of a hyaluronate or hyaluronic acid synthase gene, cDNA, and gene product (HAS), as may be used for the polymerization of glucuronic acid and N-acetylglucosamine into the glycosaminoglycan hyaluronic acid. The present invention identifies the seHAS locus and discloses the nucleic acid sequence which encodes for the enzymatically active seHAS gene from Streptococcus equisminis. The HAS gene also provides a new probe to assess the potential of bacterial specimens to produce hyaluronic acid.
[0039] Through the application of techniques and knowledge set forth herein, those of skill in the art will be able to obtain nucleic acid segments encoding the seHAS gene. As those of skill in the art will recognize, in light of the present disclosure; these advantages provide similificant utility in being able to control the expression of the seHAS gene and

control the nature of the seHAS gene product, the seHAS enzyme, that is produced.

[0040] Accordingly, the invention is directed to the isolation of a purified nucleic acid segment which has a coding region encoding enzymatically active MAS, whether it be from prokaryotic or cutsaryotic sources. This is possible because the enzyme, and indeed the gene, is one found in both eukaryotes and some prokaryotes. Eukaryotes are also known to produce HA and then have HA synthase genes that can be embloved in connection with the invention.

[0041] HA synthase-encoding nucleic acid segments of the present invention are defined as being isolated free of total chromosomal or genomic DNA such that they may be readily manipulated by recombinant DNA techniques Accordingly, as used herein, the phrase "a purified nucleic acid segment" refers to a DNA segment isolated free of unrelated chromosomal or genomic DNA and retained in a state rendering it useful for the practice of recombinant techniques, such as DNA in the form of a discrete isolated DNA fragment, or a vector (e.g., plasmid, phage or virus) incorporating such a fragment.

[0042] A preferred embodiment of the present invention is a purified nucleic acid segment having a coding region encoding enzymatically active IAS. In particular, the purified nucleic acid segment encodes the seHAS of SEQ ID NO: 2 or the purified nucleic acid segment comprises a nucleotide sequence in accordance with SEQ ID NO:1.

[0043] Another embodiment of the present invention comprises a purified nucleic acid segment having a coding region encoding enzymatically active HAS and the purified nucleic acid segment is capable of hybridizing to the nucleotide sequence of SEQ ID NO:1.

[0044] The present invention also comprises a natural or recombinant vector consisting of a plasmid, cosmid, phage, or virus vector. The recombinant vector may also comprise a purified nucleic acid segment having a coding region encoding enzymatically active IAIS.

[0045] In particular, the purified nucleic acid segment encodes the seHAS of SEQ ID NO:2 or the purified nucleic

acid segment comprises a nucleotide sequence in accordance with SEQ ID NO.1, If the recombinant vector is a plasmid, it may further comprise an expression vector. The expression vector may also include a promoter operatively linked to the enzymatically active IAS coding region.

[0046] In another preferred embodiment, the present invention comprises a recombinant host cell such as a prokaryotic cell transformed with a recombinant vector. The recombinant vector includes a purified nucleic acid segment having a coding region encoding enzymatically active HAS. In particular, the purified nucleic acid segment encodes the seHAS of SEQ ID NO.2 or the purified nucleic acid segment comprises a nucleotide sequence in accordance with SEQ ID NO.1. (0047) The present invention also comprises a recombinant host cell, such as an eukaryotic cell transfected with a recombinant vector comprising a purified-nucleic acid segment having a coding region encoding enzymatically active HAS. In particular, the purified nucleic acid segment encodes the seHAS of SEQ ID NO.2 or the purified nucleic acid segment comprises a nucleotide sequence in accordance with SEQ ID NO.1. The concept is to create a specifically modified seHAS gene that encodes an enzymatically active HAS capable of producing a hyaluronic acid polymer having a modified structure or a modified size distribution.

[0048] The present invention further comprises a recombinant host cell which is electroporated to introduce a recombinant vector into the recombinant host cell. The recombinant vector may include a purified nucleic soid segment having a coding region encoding enzymatically active HAS. In particular, the purified nucleic acid segment encodes the seHAS of SEQ ID NO.2 or the purified nucleic acid segment comprises a nucleotide sequence in accordance with SEQ ID NO.1. The enzymatically active HAS may also be capable of producing a hyaluronic acid polymer having a modified structure or a modified size distribution.

[0049] In yet another preferred embodiment, the present invention comprises a recombinant host cell which is transduced with a recombinant vector which includes a purified nucleic acid segment having a coding region encoding enzymatically active HAS. In particular, the purified nucleic acid segment encodes the seHAS of SEQ ID NO:2 or the purified nucleic acid segment comprises a nucleotide sequence in accordance with SEQ ID NO:1. The enzymatically active HAS is also capable of producing a hyaluronic acid polymer having a modified structure or a modified size distribution.

[0850] The present invention also comprises a purified composition, wherein the purified composition comprises a polypeptide having a coding region encoding enzymatically active HAS and further having an amino acid sequence in accordation of this SEQ ID NO 2.

[0051] In another embodiment, the invention comprises a method for, detecting a DNA species, comprising the steps of . (1) obtaining a DNA semple. (2) contacting the DNA sample with a purified nucleic acid segment in accordance with SEQ ID NO 1; (3) hybridizing the DNA sample and the purified nucleic acid segment thereby forming a hybridized comblex, and (4) detection the complex.

[0052] The present invention also comprises a method for detecting a bacterial cell that expresses mRNA encoding seHAS, comprising the steps of: (1) obtaining a bacterial cell sample; (2) contacting at least one nucleic acid from the bacterial cell sample with purified nucleic acid segment in accordance with SEQ ID Not1; (3) hydridizing the stast one nucleic acid and the purified nucleic acid segment thereby forming a hybridized complex; and (4) detecting the hybridized complex, wherein the presence of the hybridized complex is indicative of a bacterial strain that expresses mRNA encoding seHAS.

[0053] The present invention also comprises methods for detecting the presence of either seHAS or spHAS in a cell.
In particular, the method comprises using the oligonucleotides set forth in Seq. [10 Nos: 3-3 as probes. These oligonucleotides would a allow a practitioner to search and detect the presence of SeHAS or spHAS in a cell.

10054] The present invention further comprises a method for producing hyaluronic acid, comprising the steps of. (1) introducing a purified nucleic acid segment having a coding region encoding enzymatically active HAS into a bost organism, wherein the host organism contains nucleic acid segments encoding enzymes which produce UDP-GloNAc and UDP-GloX, (2) growing the host organism in a medium to secrete hyaluronic acid; and (3) recovering the secreted hyaluronic acid.

[0055] The method may also include the step of extracting the secreted hyalutonic acid from the medium as well as the step of purifying the extracted hyalutonic acid. Furthermore, the host organism may secrete a structurally modified hyalutonic acid or a size modified hyalutonic acid.

[0056] The present invention further comprises a pharmaceutical composition comprising a preselected pharmaceutical drug and an effective amount of hyaluronic acid produced by a recombinant HAS. The pharmaceutical composition may have a hyaluronic acid having a modified molecular weight pharmaceutical composition capable of evading an immune response. The modified molecular weight may also produce a pharmaceutical composition capable of targeting a specific tissue or cell type within the patient having an affinity for the modified molecular weight pharmaceutical composition.

[0057] The present invention also comprises a purified and isolated nucleic acid sequence encoding enzymatically active seHAS, where the nucleic acid sequence is (a) the nucleic acid sequence in accordance with SEQ ID NO.1: (D) complementary nucleic acid sequences has occordance with SEQ ID NO.1: (D) in fuciency nucleic acid sequences have sequence in accordance with SEQ ID NO.1: (D) in fuciency nucleic acid.

acid sequences which will hybridize to the nucleic acid in accordance with SEQ ID NO:1; and (d) nucleic acid sequences which will hybridize to the complementary nucleic acid sequences of SEQ ID NO:1.

[0058] The present invention further comprises a purified and isolated nucleic acid segment consisting essentially of a wucleic acid segment encoding enzymatically active HAS.

- [0059] The present invention also comprises an isolated nucleic acid segment consisting essentially of a nucleic acid segment encoding setHAS having a nucleic acid segment sufficiently duplicative of the nucleic acid segment in accordance of SEQ ID NO.1 to allow possession of the biological property of encoding for an enzymatically active HAS. The nucleic acid segment may also be a cDNA sequence.
- [0060] The present invention also comprises a purified nucleic acid segment having a coding region encoding enzymatically active IAS, wherein the purified nucleic acid segment is capable of hybridizing to the nucleotide sequence in accordance with SEO ID No.1.

BRIEF DESCRIPTION OF THE SEVERAL VIEWS OF THE DRAWINGS

15 [0061]

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- FIG. 1 depicts that cross hybridization between seHAS and spHAS genes does not occur.
- FIG. 2 figuratively depicts the relatedness of seHAS to the bacterial and eukaryotic HAS proteins.
- FIG. 3 figuratively depicts evolutionary relationships among some of the known hyaluronan synthases.
- FIG. 4 depicts the HA size distribution produced by various engineered Streptococcal HAS enzymes.
- FIG. 5 figuratively depicts the overexpression of recombinant seHAS and spHAS in E. coli.
- FIG. 6 depicts purification of Streptococcal HA synthase.
- FIG. 7 depicts a gel filtration analysis of HA synthesized by recombinant streptococcal HAS expressed in yeast membranes.
- FIG. 8 is a Western blot analysis of recombinant seHAS using specific antibodies.
 - FIG. 9 is a kinetic analysis of the HA size distributions produced by recombinant seHAS and spHAS.
 - FIG. 10 graphically depicts the hydropathy plots for seHAS and predicted membrane associated regions.
 - FIG. 11 is a graphical model for the topologic organization of seHAS in the membrane.
 - FIG. 12 is a demonstration of the synthesis of authentic HA by the recombinant seHAS.
- FIG. 13 depicts the recognition of nucleic acid sequences encoding seHAS, encoding spHAS, or encoding both seHAS and spHAS using specific oligonucleotides and PCR.
- FIG. 14 depicts aligonucleatides used for specific PCR hybridization.

DETAILED DESCRIPTION OF THE INVENTION

[0062] Before explaining at least one embodiment of the invention in datal, it is to be understood that the invention is not limited in its application to the details of construction and the arrangement of the components set forth in the following description or illustrated in the drawings. The invention is capable of other embodiments or of being practiced or carried but in various ways. Also, it is to be understood that the phraseology and terminology employed herein is for purpose of description and should not be recarded as limiting.

[9063] As used herein, the term "nucleic acid segment" and "DNA segment" are used interchangeably and refer to a DNA onelocule which has been isolated free of total genomic DNA of a particular species. Therefore, a "purified" DNA or nucleic acid segment as used herein, refers to a DNA segment which contains a Hyaluronate Synthase ("HAS") coding sequence yet it is loalted away from, or purified free from, unrelated genomic DNA, for example, including coccus equismilla or, for example, mammalian host genomic DNA. Included within the term "DNA segment", and has generals and smaller fragments of such segments, and also recombinant vectors, including, for example, plasmids, cosmids, shape, viruses, and the like.

[0044] Similarly, a DNA segment comprising an isolated or purified setNAS gene refers to a DNA segment including HAS coding sequences is latited substantially wave from other naturally occurring genes or protein encoding sequences. In this respect, the term "gene" is used for simplicity to refer to a functional protein, polypeptide or peptide encoding unit. As will be understood by those in the art, this functional term includes genomic sequences. CDNA sequences or combinations thereof. "Isolated substantially away from other coding sequences" means that the gene of interest, in this case setNAS, forms the significant part of the coding region of the DNA segment, and that the DNA segment does not contain large portions of naturally-occurring coding DNA, such as large chromosomal fragments or other functional genes or DNA coding regions. Of course, this refers to the DNA segment as originally isolated, and does not exclude genes or coding regions.

[0065] Due to certain advantages associated with the use of prokaryotic sources, one will likely realize the most advantages upon isolation of the HAS gene from prokaryotes such as S. pyogenes, S. equisimilis, or P. multocida. One

such advantage is that. typically, eukaryotic enzymes may require significant posttranslational modifications that can only be enthered in a eukaryotic host. This will tend to limit the applicability of any eukaryotic HA, synthase gene that is obtained. Moreover, those of ordinary skill in the art will likely realize additional advantages in terms of time and ease of genetic manipulation where a prokaryotic enzyme gene is sought to be employed. These additional advantages include (a) the sease of isolation of a prokaryotic gene because of the relatively small size of the genome and, therefore, the reduced amount of screening of the corresponding genomic library and (b) the ease of manipulation because the overall size of the odding region of a prokaryotic gene is significantly smalled due to the absence of introns. Furthermore, if the product of the seHAS gene (i.e., the enzyme) requires posttranslational modifications, these would best be achieved in a similar profavorious cellular environment (host) from which the quere was derived.

[0066] Preferably, DNA sequences in accordance with the present invention will further include genetic control regions which allow the expression of the sequence in a selected recombinant host. Of course, the nature of the control region employed will generally vary depending on the particular use (e.g., cloning host) envisioned.

[0067] In particular embodiments, the invention concerns isolated DNA segments and recombinant vectors morporating DNA sequences which encode a seHAS gene, that includes within its amino acid sequence an amino acid sequence in accordance with SEO ID NO.2. Moreover, in other particular embodiments, the invention concerns isolated DNA segments and recombinant vectors incorporating DNA sequences which encode a gene that includes within its amino acid sequence of an HAS gene or DNA, and in particular to an HAS gene or CNA, corresponding to Streptococcus equisimilis HAS. For example, where the DNA segment or vector encodes a full length HAS protein, or is intended for use in expressing the HAS protein, preferred sequences are those which are essentially as set fort in SEO ID NO.2.

[0068] Nucleic acid segments having HA synthase activity may be isolated by the methods described herein. The term "a sequence essentially as set forth in SEQ ID NO.2" means that the sequence substantially corresponds to a portion of SEQ ID NO.2 and has relatively free marino acids which are not defented to, or a biologically functional equivalent of, the amino acids of SEQ ID NO: 2. The term "biologically functional equivalent" is well understood in the art and is further defined in detail herein, as a gene having a sequence essentially as set forth in SEQ ID NO:2, and that is associated with the shifty of prokaryotes to produce HA or a hyaluronia caic doat.

[0069] For instance, the seHAS and spHAS coding sequences are approximately 70% identical and rich in the bases adenine (A) and thymine (T). SeHAS base content is A.26 T/%. C-19 13%, C-20.31 %, and T-33.3% (AT = 60%). Whereas spHAS is A-31.34%, C-18.23%, G-18.34%, and T-35.8% (AT = 67%). Those of ordinary skill in the art would be surprised that the seHAS coding sequence does not hybridize outlid be due to short interruptions of mismatched bases throughout the open reading frames. The inability of spHAS and seHAS to cross-hybridize is shown in FIG. 1. The longest stretch of identical inuicatedise common to both the seHAS and the spHAS coding sequences is not 20 nucleotides. In addition, the very A-T rich sequences will form less stable hybridization complexes than G-C rich sequences. Another possible explanation could be that there are several stretches of As or Ts in both sequences that could hybridize in a misaligned and unstable manner. This would put the seHAS and spHAS gene sequences out of frame with respect to each other, thereby decreasing the probability of productive hybridization.

[0070] Because of this unique phenomena of two genes encoding proteins which are 70% identical not being capable of cross-hybridizing to one another, it is beneficial to think of the claimed nucleic acid segment in terms of its function, i.e. a nucleic acid segment which encodes enzymatically active hyaturonate synthase. One of ordinary skill in the art would appreciate that a nucleic acid segment encoding enzymatically active hyaturonate synthase may contain conserved or semi-conserved substitutions to the sequences set forth in SEQ ID NOS: 1 and 2 and yet still be within the scope of the invention.

[6071] In particular, the art is replete with examples of practitioners ability to make structural changes to a nucleic acid segment (i.e. encoding conserved or semi-conserved amino acid substitutions) and still preserve its enzymatic or functional activity. See for example: (1) Risler et al. "Amino Acid Substitutions in Structurally Related Proteins. A Pattern Recognition Approach". J. Mol. Biol. 204.1019-1029 (1988) [*]... according to the observed exchangeability of amino acid side chains, only four groups could be delineated; (i) the and Val; (ii) Lou and Met, (iii) Lys. Arg, and Gin, and (iv) Yr and Phe."]. (2) Niefind et al. "Amino Acid Similarity Coefficients for Protein Modeling and Sequence Alignment. Derived from Main-Chain Folding Anoles "J. Mol. Biol. 2148.1-497 (1991) (similarity parameters allow amino acid substitutions to be designed; and (3) Overington et al. "Environment-Specific Amino Acid Substitution Tables: Tertiary Templates and Prediction of Protein Foldies, "Protein Science 1:16-226 (1992) (Phalysles) of the pattern of observed substitutions as a function of local environment shows that there are distinct patterns..." Compatible changes can be made.]

[0072] These references and countiess others, indicate that one of ordinary skill in the art, given a nucleic acid sequence, could make substitutions and changes to the nucleic acid sequence without changing its functionality. Also, a substituted nucleic acid segment may be highly identical and retain its enzymatic activity with regard to its unadulterated parent and vet still field to hybridize thereto.

[0073] The invention discloses nucleic acid segments encoding enzymatically active hyaluronate synthase - seHAS and spHAS. Although seHAS and spHAS are 70% identical and both encode enzymatically active hyaluronate synthase, they do not cross hybridize. Thus, one of ordinary skill in the art would appreciate that substitutions can be extended to the seHAS nucleic acid segment listed in SEQ ID NO. 1 without deviating outside the scope and claims of the present invention. Standardized and accepted functionally equivalent armino acid substitutions are presented in Table 1.

TABLE

	77 500 50.00
Amino Acid Group	Conservative and Semi-Conservative Substitutions
NonPolar R Groups	Alanine, Valine, Leucine, Isoleucine, Proline, Methionine, Phenylalanine, Tryptophan
Polar, but uncharged, R Groups	Glycine, Serine, Threonine, Cysteine, Asparagine, Glutamine
Negatively Charged R Groups	Aspartic Acid, Glutamic Acid
Positively Charged R Groups	Lysine, Arginine, Histidine

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[0074] Another preferred embodiment of the present invention is a purified nucleic acid segment that encodes a protein in accordance with SEQ ID NO: 2, further defined as a recombinant vector. As used herein, the term "recombinant vector" refers to a vector that has been modified to contain a nucleic acid segment that encodes an HAS protein, or fragment thereof. The recombinant vector may be further defined as an expression vector comprising a promoter operatively linked to said HAS encoding nucleic acid segment.

[0075] A further preferred embodiment of the present invention is a host cell, made recombinant with a recombinant vector comprising an HAS gene. The preferred recombinant hostcell may be a prokaryotic cell. In another embodiment, the recombinant host cell is a eukaryotic cell. As used heren, the term "engineered" or "recombinant" cell is intended to refer to a cell into which a recombinant gene, such as a gene encoding HAS, has been introduced. Therefore, engineered cells are distinguishable from naturally occurring cells which do not contain a recombinantly introduced gene. Engineered cells are thus cells having a gene or genes introduced through the hand of man. Recombinantly introduced genes will either be in the form of a cDNA gene, a copy of a genomic gene, or will include genes positioned adiacent to a romoter profit harturally associated with the particular introduced and promoter profit harturally associated with the particular introduced profits.

[0076] Where one desires to use a host other than Streptococcus, as may be used to produce recombinant HA synthase, it may be advantageous to employ a prokaryotic system such as £ coli, 8 subfilis, £ectococcus sp, or even eukaryotic systems such as yeast or Chinese hamster ovary, African green monkey kidney cells, VERO cells, or the like. Of course, where this is undertaken it will generally be desirable to bring the HA synthase gene under the control of sequences which are functional in the selected alternative host. The appropriate DNA control sequences, as well as their construction and use, are generally well known in the art as discussed in more detail hereinbelow.

[0077] In preferred embodiments, the HA synthase-encoding DNA segments further include DNA sequences, known in the art. functionally as origins of replication or "replicanes", which allow replication of configuous sequences by the particular host. Such origins allow the preparation of extrachromosomally localized, and replicating chimeric segments or plasmids, to which HA synthase DNA sequences are ligited. In more preferred instances, the employed origin is one capable of replication in bacterial hosts suitable for biotechnology applications. However, for more versatified for cloned DNA segments, it may be desirable to alternatively or even additionally employ origins recognized by other host systems whose use is contemplated (such as in a shuttle vector.)

[0078] The isolation and use of other replication origins such as the SV40, polyoma or bovine papilloma virus origins, which may be employed for cloning or expression in a number of higher organisms, are well known to those of ordinary skill in the art. In certain embodiments, the invention may thus be defined in terms of a recombinant transformation vector which includes the HA synthase coding gene sequence together with an appropriate replication origin and under the control of selected control regions.

[0079] Thus, it will be appreciated by those of skill in the art that other means may be used to obtain the HAS geneor cDNA, in light of the present disclosure. For example, polymerase chain reaction or RT-PCR produced DNA Sprements may be obtained which contain full complements of genes or cDNAs from a number of sources, including other strains of Streptococcus of from eukaryotic sources, such as cDNA libraries. Virtually any molecular cloning approach may be employed for the generation of DNA fragments in accordance with the present invention. Thus, the only limitation generally on the particular method employed for DNA isolation is that the isolated nucleic acids should encode a biologicality functional equivalent HA synthasy.

[0880] Once the DNA has been isolated it is ligated together with a selected vector. Virtually any cloning vector can be employed to realize advantages in accordance with the invention. Typical useful vectors include plasmids and phages for use in protaryotic organisms and even viral vectors for use in eukaryotic organisms. Examples include

pKK223-3, pSA3, recombinant lambda. SV40, polyoma, adenovirus, bovine papilloma virus and retroviruses. However, it is believed that particular advantages will ultimately be realized where vectors capable of replication in both *Lacto-coccus of Bacillus* stains and *L. coli* are employed.

[0081] Vectors such as these, exemplified by the pSA3 vector of Dao and Ferretti or the pAT19 vector of Tineu-Cuct, et al., allow one to perform clonal colony selection in an easily manipulated host such as E. Colf, followed by subsequent transfer back into a food grade Lacfococcus or Bacillus strain for production of HA. These are benign and well studied organisms used in the production of certain floods, and biotechnology products. These are advantageous in that one can augment the Lacfococcus or Bacillus strain's ability to synthesize HA through gene dosaging (i.e., providing extra copies of the HA synthase gene by amplification) and/or inclusion of additional genes to increase the availability of HA procursors. The inherent ability of a bacterium to synthesize HA can also be augmented through the formation of at copies, or amplification, of the plasmed that carries the HA synthase gene. This amplification can account for up to a 10-fold increase in plasmid, copy number and therefore, the HA synthase gene copy number.

10-fold increase in plasmid, copy number and, therefore, the HA synthase gene copy number.

[00s2] Another procedure that would further augment HA synthase gene copy number is the insertion of multiple copies of the gene into the plasmid. Another technique would include integrating the HAS gene into chromosomal DNA. This extra amplification would be especially leasible, since the bacterial HA synthase gene size is small. In owns scenarios, the chromosomal DNA-ligated vector is employed to transfect the host that is selected for clonal screening purposes such as E.-Co./I through the use of a vector that is capable of expressing the inserted DNA in the chosen host ideals to proceed initially by preparting a cDNA library. This is carried out first by isolation of mRNA from the above cells, followed by preparation of double stranded cDNA using an enzyme with reverse transcriptise activity and ligitation with the selected vector Numerous possibilities are available and known in the art for the preparation of the double stranded cDNA, and all such techniques are believed to be applicable. A preferred technique involves reverse transcriptered techniques involves reverse transcriptered techniques involves reverse transcriptered techniques, such as by ligation into the appropriate vector and amplification in the appropriate host by accepted techniques, such as by ligation into the appropriate vector and amplification in the appropriate host by the techniques set forth herein, one may desire to employ phage expression vectors, such as £gt11, £g112, £Gem11, and/or £APA for the cloning and expression screening of cDNA clones as forther inclining and coloning and expression screening of cDNA clones and coloning and expression screening of cDNA clones.

[0084] In certain other embodiments, the invention concerns isolated DNA segments and recombinant vectors that include within their sequence an uncleic acid sequence essentially as set forth in SEQ ID-NO.1 The term "essentially as set forth in SEQ ID-NO.1" his used in the same sense as described above and means that the nucleic acid sequence substantially corresponds to a portion of SEQ ID ND;1, and has relatively few codons which are not identical, or functionally equivalent, to the codons of SEQ ID NO.1. The term "incritionally equivalent codon" is used herein to refer to codons that encode the same amino acid, such as the six codons for arginine or serine, as set forth in Table I, and also refers to codons that encode biologically equivalent amino acids.

[0055] It will also be understood that amino acid and nucleic acid sequences may include additional residues, such as additional N-or C-terminal amino acids or 5' or 3' nucleic acid sequences, and yet still be essentially as set forth in one of the sequences disclosed herein, so long as the sequence meets the criteria set forth above, including the maintenance of biological protein activity where protein expression and enzyme activity is concerned. The addition of terminal sequences particularly applies to nucleic acid sequences which may, for example, include various non-coding sequences flanking either of the 5' or 3' portions of the coding region or my include various internal sequences, which are known to occur within genes. In particular, the amino acid sequence of the HAS gene in eukaryotes appears to be 40% larger than that four in prokarvotes.

[0085] Allowing for the degeneracy of the genetic code as well as conserved and semi-conserved substitutions, sequences which have between about 40% and about 90%, or even more preferably, between about 80% and about 90%, or even more preferably, between about 90% and about 90% of nucleotides which are identical to the nucleotides of SEC ID NO.1 will be sequences which are "essentially as set forth in SEC ID NO.1". Sequences which are essentially the same as those set forth in SEC ID NO.1 may also be functionally defined as sequences which are capable of hybridizing to a nucleic acid segment containing the complement of SEC ID NO.1 under standard or less stringent hybridizing conditions. Suitable standard hybridization conditions will be well known to those of skill in the art and are clearly set forth herein.

[0087] The term "standard hybridization conditions" as used herein, is used to describe those conditions under which substantially complementary uncleic acid segments will form standard Watson-Chick base-paining. A number of factors are known that determine the specificity of binding or hybridization, such as pH, temperature, salt concentration, the presence of agents, such as formamide and dimethyl sulfoxide, the length of the segments that are hybridizing, and the like When it is contemplated that shorter nucleic acid segments will be used for hybridization, for example fragments between about 14 and about 100 nucleotides, salt and temperature preferred conditions for hybridization will include 1.2-1.8 x HPS at 40-50°C.

[0088] Naturally, the present invention also encompasses-DNA segments which are complementary, or essentially

complementary, to the sequence set forth in SEQ ID NO.1. Nucleic acid sequences which are "complementary" if are those which are capable of base-pairing according to the standard Watson-Crick complementarity rules. As used herein, the term "complementary sequences" means nucleic acid sequences which are substantially complementary, as may be assessed by the same nucleotide comparison set forth above, or as defined as being capable of hybridizing to the nucleic acid sequences of SEQ ID NO.1

[0089] The nucleic acid segments of the present invention, regardless of the length of the coding sequence itself, may be combined with other DIM-sequences, such as promoters, polyadenylation signals, additional restriction enzyme sites, multiple doming sites, epitope tags, poly histidine regions, other coding segments, and the like, such that overall length may vary considerably. It is therefore contemplated that a nucleic acid fragment of almost any length may be employed, with like total length preferably being limited by the ease of preparation and use in the intended recombinant DIM protocol.

[0990] Naturally, it will also be understood that this invention is not limited to the particular nucleic acid and amino acid sequences of SEQ ID NO: 1 and 2. Recombinant vectors and isolated DNA segments may therefore variously include the IHAS coding regions themselves, coding regions bearing selected alterations or modifications in the basic coding region, or they may encode larger polypeptides which nevertheless include IHAS-coding regions or may encode blookpically functional equivalent proteins or peoplides which have variant amino acids sequences.

[0081] For instance, we have found, characterized, and purified hyalutonate synthase in two other systems: (a) the gram-negative bacteria Pasturella multiocida (SEQ ID NO:19); and (2) chlorella rirus PBCV-1 (SEQ ID NO:87 and 8). The presence of hyalutonan, ynthase in these two systems and our ability to purify and use the hyalutonan synthase from these two different systems indicates our ability to purify and isolate nucleic acid sequences encoding enzymatically active hyalutonan synthase.

[0092] The capsule of Carter Type A P multocide (SEQ ID NO:19) was long suspected of containing hyaluronic acid-HA. characterization of the HA synthase of P multocida led to interesting enzymological differences between it and the selfAS and soHAS proteins.

[0093] P. multicolds cells produce a readily visible extracellular HA capsule, and since the two streptococcal HASs are membrane proteins, membrane prosperations of the fowl cholers pathogen were tested. In early trials, oruse membrane fractions derived from ultrasonication alone possessed very low levels of UDP-GlicN-capendent UDP- $\frac{1}{1}$ Cl GlicA incorporation into HA[-0.2 pmol of GlicA transfer (iii) of proteins) $^{-1}$ + $^{-1}$ when assayed under conditions similar to those for measuring streptococcal HAS activity. The enzyme from E. con with the recombinant hash plasmid was also recabilitant to isolation at first. These results were in contrast to the easily detectable amounts obtained from Strepto-

coocus by similar methods. [004] An alternative preparation protocol using ice-cold lysozyme treatment in the presence of protease inhibitors in conjunction with ultrasonication allowed the substantial recovery of HAS activity from both species of Gram-negative bacteria. Specific activities for HAS of 5-10 pinol of Glick transferred (ug of protein): "h¹- vare routinely obtained for crude membranes of wild-type P: multocide with the new method. In the absence of UDP-GlicNac, virtually no radioactivity (<1% of identical assay with both sugar precursors) from UDP-[1**C]olicA was incorporated into higher molecular weight material. Membranes repeared from the acapsular mutant, ThA, possessed no detectable HAS activity when supplemented with both sugar nucleotide precursors (data not shown). Gel-fitration analysis using a Sephacryl S-200 oclumn indicates that the molecular mass of the majority of the "C-labeled product synthesized in virtie is 28 x 10.4 Da since the material elutes in the void volumes, such a -value corresponds to a HA molecule composed of at least 400 monomers. This product is servisitive to Streptomores hylumonidase diagestion but resistant to protease treatment.

[0095] The parameters of the HAS assay were varied to maximize incorporation of UDP-sugars into polysaccharide by P. multocida membranes. Steptococcal spHAS requires Mg²⁺ and therefore this metal ion was included in the initial assays of P. multocida membranes. The P. multocida HAS (pmHAS) was relatively active from pH 6.5 to 8.6 in Tristype buffers with an optimum at pH 7. The HAS activity was linear with respect to the incubation time at neutral pH for at least 1 h. The pmHAS was apparently less active at higher ionic strengths because the addition of 100 mM NaCl to the reaction containing 5.0 mM Tris, pH 7, and 2 mM MaCl. The educed square incorporation by ~60%.

[0096] The metal ion specificity of the pmHAS was assessed at pH 7. Under metal-free conditions in the presence of EDTA, no incorporation of radiolabeled precursor into physaccharide was detectable (<0.50 of maximal signal) Mn²⁺ gave the highest incorporation rates at the lowest ion concentrations for the tested metals (Mg, Mn, Co, Cu, and Ni). Mg²⁺ gave about 00% of the Mn²⁺ simulation but at 10-fold higher concentrations. Co²⁺ or N²⁺ at 10 mM supported lower levels of activity (20% or 9%, respectively, of 1 mM Mn²⁺ assays), but membranes supplied with 10 mM Cu²⁺ were inactive. Indeed, mixing 10 mM Cu²⁺ and 20 mM²⁺ Mg²⁺ with the membrane preparation resulted in almost no incorporation of label into polysaccharials (<0.3% of Mg only value).

[0097] Initial characterization of the pmHAS was performed in the presence of Mg²⁺. The binding affinity of the enzyme for its sugar nucleotide precursors was assessed by measuring the apparent K_M, value. Incorporation of [¹⁴C] GloA or [⁹H]GloNAc into polysaccharide was monitored at varied concentrations of UDP-GloNAc or UDP-GloA, respectively. In Mg²⁺-containing buffers, the apparent K_M, values of -20 µM for UDP-GloA and -75 µM for UDP-GloA vere

determined utilizing Hanes-Woolf plots ([S]/v versus [S]) of the titration data. The $V_{\rm max}$ values for both sugars were the same because the slopes, corresponding to $1/V_{\rm max}$, of the Hanes-Woolf plots were equivalent. In comparison to results from assays with $M_{\rm p}^{2+}$, the $K_{\rm hx}$ value for UDP-GicNAc was increased by about 25-50% to -105 μ M and the $V_{\rm max}$ increased by a factor of 2-3-fold in the presence of $M_{\rm p}^{2+}$.

10088] The HA synthase enzymes from either P multicide. S. equisimitis, or S, progenes utilizes UDP-sugars, but they possess somewhat different kinetic optima with respect to pH and metal ion dependence and $K_{\rm M}$ values. The enzymes are most active at $y_{\rm H}$? however, the pmHAS reportedly displays more activity at slightly actic pH and is relatively inactive above pH 7.4. The pmHAS utilizes Mn^2 more efficiently than Mg^2 : under the m with assignment of the physicological metal cofactor in the bacterial cell is unknown. In comparison, in previous studies with the streptococcal enzyme, Mg^2 : was much better than Mn^{2k} but the albeit smaller effect of Mn^{2k} : was maximal at -10-dold lower concentrations than the optimal Mg^{2k} concentration. The pmHAS apparently binds the UDP-sugars more tightly than spHAS. The measured $K_{\rm M}$ values for the pmHAS in crude membranes are about 2-3-dold lower for each substrate than those obtained from the HAS found in streptococcal membranes. 50 or 39 μ M for UDP-GlcA and 500 or 150 μ M for UDP-GlcA, respectively.

[0099] By kinetic analyses, the V_{max} of the pmHAS was 2-3-fold higher in the presence of Mn²⁺ than Mg²⁺, but the UDP-ClicNAC K_M value was increased slightly in assays with the former ion. This observation of epparent lowered affinity suggests that the increased polymerization rate was not due to better fulning of the Mn²⁺ ion/sugar nucleotide complex to the enzyme active site(s). Therefore, it is possible that Mn²⁺ enhances some other reaction step, alters another site/structure of the enzyme, or modifies the phospholipid membrane environment. The gene sequence and the protein sequence of pmHAS are shown in SEQ ID NO-19.

[0100] Chlorella virus PBCV-1 encodes a functional glycosyltransferase that can synthesize a polysaccharide, hyaluronan hysturonic acid, HA]. This finding is contrary to the general observation that viruses either; (a) utilize host onel glycosyltransferases to create new carbohydrate structures, or (b) accumulate host cell glycosyltransferases) to create new carbohydrate structures, or (b) accumulate host cell glycocorpiugates that principle of the property of the propert

[0101] The vertebrate HAS enzymes (DG42, HAS1, HAS2, HAS3) and streptococcal HasA enzymes (spHAS and seHAS) have several regions of sequence similarity. While sequencing the doublestranded DNA genome of virus PB-CV-1 [Parametoim pursair exhortes] and ORF [open reading frame]. ASR (Accession #442580), encoding a 567 residue protein with 28 to 33% amino acid identity to the various HASs was discovered. This protein is designated cvHAS (chlorella virus HA synthase). The gene sequence encoding PBCV-1 and its protein sequence are shown in SEQ ID NOS/7 and 8.

SECULIVIOS.7 and 6.
[1012] PSCV-1 is the prototype of a family (Phycodnarviridae) of large (175-130 nm diameter) polyhedral, plaqueforming viruses that replicate in certain unicellular, eukaryotic chlorella-like green algae. PSCV-1 virions contain at least 50 different proteins and a lipid component locateld inside the outer glycoptoien capsid. The PSCV-1 genome is

a linear, nonpermuted 330-kb dsDNA molecule with covalently closed hairpin ends.

[9103] Based on its deduced amino acid sequence, the A98R gene product should be an integral membrane protein. To test this hypothesis, recombinant A98R was produced in Ecohernicis cold and the membrane fraction was assigned for HAS activity. UDP-GIcA and UDP-GIcNAc were incorporated into the polysaccharide by the membrane fraction derived from cells containing the A98R gane on a plasmid, pCVHAS, (average specific activity 2.5 pmoles Glock transferring protein/min), No activity as detected in the solubile fraction of cells transformed with pCVHAS, UDP-GIcA and UDP-GIcNAc were simultaneously required for polymerization. The activity was optimal in Neges buffer apth 7.2 in the presence of 10 mM MrCl₂, whereas no activity was detected if the metal ion was omitted. Mg²⁺ and CO²⁺ were -20% as effective as Mm²⁺ at similar concentrations. The pmHAS has a similar metal requirement, but other HASs prefer Mg²⁺.

[0104] The recombinant A98R enzyme synthesized a polysacoharide with an average molecular weight of 3-6xt 10⁸ Da which is smaller than that of the HA synthesized by recombinant sph4S or D642 xHAS in vito (-10⁷ Da and -5-9xt)0¹⁸ Da, respectively; 13,15). The polysacoharide was completely degraded by *Streptomyces hyalinonitics* HA lyase, an enzyme that depolymerizes HA, but not structurally related glycosanihoglycans such as heparin and chondrolin.

[0105] PBCV-1 infected chlorella cells were examined for A98R gene expression. A -1,700-nucleotide A98R transcript appeared at ~15 min post-infection and disappeared by 60 min after infection indicating that A98R is an early gene. Consequently, membrane fractions from uniffected and PBCV-1 infected chlorella cells were assayed at 50 and 90 min post-infection for HAS activity, Infected cells, but not uninfected cells, but add activity. Like the bacterially derived recombinant A98R enzyme, radiolabel incorporation from UDP-1/4°C [Glok Into pylasecharide depended on both Mark and UDP-GlokAc. This radiolabeled produce was also degraded by HA lyase. Disrupted PBCV-1 virions had no HAS

[0106] PBCV-1 infected chlorella cells were analyzed for HA polysaccharide using a highly specific 125Habeled HA-

binding protein. Extracts from cells at 50 and 90 min post-infection contained substantial amounts of HA, but, not extracts from uninfected algae or dissuped PBCV4 virions. The labeled HA-binding protein also interacted with intact infected cells at 50 and 90 min post-infection, but not healthy cells. Therefore, a considerable portion of the newly synthesized HA polysacchande was immobilized at the outer cell surface of the infected algae. The extracellular HA does not play any obvious role in the interaction between the virus and its algal host because neither plaque size or praque member was altered by including either testicular hyaluronidase (465 units/ml) or free HA polysaccharide (100 µg/ml) in the top agar of the PBCV1 losious assay.

[0107] The PBCV-1 genome also has additional genes that encode for an UDP-Glc dehydrogenase (UDP-Glc DH) and a glutamine fructose-6-phosphate aminotransferase (GRFI). UDP-Glc DH converts UDP-Glc into UDP-GlcA, a required precursor for HA biosynthesis GRFI converts fructose-6-phosphate, an interpretate in the UDP-GlcNAc metabolic pathway. Both of these PBCV-1 genes, like the A98R HAS, are expressed early in infection and encode enzymatically active proteins. The presence of multiple enzymes in the HA biosynthesis pathway indicates that HA production must serve an important function in the file cycle of the chlorella viruses.

[0108] HA synthases of Straptococcus, vertebrates, and PBCV-1 pessess many motifs of 2 to 4 residues that occur in the same relative order. These conserved moits probably reflect domains crucial for HA biosynthesis as shown in FIG. 2. The protein sequences of Group C sehAS, Group A spHAS, murine HAS1, HAS2, HAS3, and frog HAS are shown aligned in FIG. 2. The alignment of FIG. 2 was accomplished using the DNAss multiple alignment program. Residues in seHAS identified in other known HAS family members (including human HAS1 and 2, not shown) are denoted by shading and seterisks. The amino acids inclicated by dots are conserved in all members of the larger [5]-glycosyl transferses family. The claimond symbol indicates the highly conserved cysteine residue that may be crucial for enzyme activity. The approximate mid-points of predicted membrane domains MD1 through MD7 are indicated with arrows. X1 indicates X evolus Servis, and MM denotes Mue mysculis.

[0109] Regions of similarity between HASs and other enzymes that synthesize 8-linked polysacoharides from UDPsugar precursors are also being discovered as more glycosythansferases are sequenced. Examples include bacterial cellulose synthase, fungal and bacterial chitin synthises, and the various HASs. The significance of these similar structural motifs will become more apparent as the three-dimensional structures of glycosyftransferases accumulate. [0110] FIG. 3 depicts the evolutionary relationships among the from hyphomena synthase. The phylogenetic red of FIG. 3 was generated by the Higgins-Sharp algorithm using the DNAsis multiple alignment program. The calculated matching percentages are included at each branch of the dendrogram.

10 [0111] The DNA segments of the present invention encompass biologically functional equivalent HAS proteins and peptides. Such sequences may arise as a consequence of codon redundancy and functional equivalency which are known to occur naturally within nucleic acid sequences and the proteins thus encoded. Alternatively, functionally equivalent proteins or peptides may be created via the application of recombinant DNA (achnology, in which changes in the protein structure may be engineered, based on considerations of the properties of the amino acids being exchanged. Changes designed by man may be introduced through the application of site-directed mutagenesis techniques, e.g., to introduce improvements to the enzyma activity or to antigenicity of the HAS protein or to test HAS mutants in order to examine HA synthase activity at the molecular level.

[0112] Also, specific changes to the HAS coding sequence can result in the production of HA having a modified size distribution or structural configuration. One of ordinary skill in the art would appreciate that the HAS coding sequence can be manipulated in a manner to produce an altered hyaluronate synthase which in turn is capable of producing hyaluronic acid having differing polymer sizes ander functional capabilities. For example, the HAS coding sequence may be altered in such a manner that the hyaluronica synthase has an altered sugar substrate specificity so that the hyaluronicate synthase creates a new hyaluronicacid-like polymer incorporating a different structure such as a previously unincorporated sugar or sugar derivality. This newly incorporated sugar could result in a modified hyaluronic acid having a smaller or larger polymer size/molecular weight, or both. As will be appreciated by one of ordinary skill in the art given the HAS coding sequences, changes and/or substitutions can be made to the HAS coding sequences such that these desired property and/or size modifications can be accomplished.

TABLE

	IMPLE II								
Sugar nucleotide specifi	city and Magnesium ion requiremen	t of recombinant seHAS							
	HA Synthesis*								
Second Sugar nucleotide present (μΜ)	UDP-[1ªC]GlcA dpm (%)	UDP- [3H] GlcNAc dpm (%)							
None	90 (2.1%)	8 (1.2%)							
UDP-GloNAc (300)	4134 (100%)	******							
UDP-GlcA C120)	*****	635 (100%)							
UDP-Glc (160)	81 (1.9%)	10 (1.5%)							
UDP-GaINAc (280)	74 (1.7%)	19 (2.9%)							
UDP-GalA (150)	58 (1.4%)	19 (2 9%)							
UDP-GIENAC + EDTA	31 (0.7%)								
UDP-GlcA + EDTA		22 (3.4%)							

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* Membranes (324 ng protein) were incubated at 37°C for th with either 120 µM UDP. [14C] GloA (2 8x fd⁴ dpm) or 300 µM UDP. [14H]GloRAc (2x fd⁴ dpm). The radiobabeds sugar nucleotide was steed in the presence of the inacisted second nonlabeled sugar nucleotide. HA synthase activity was determined as described in the insubsection.

[0113] The term "modified structure" as used herein denotes a hyaluronic acid polymer containing a sugar or derivtative not normally found in the naturally occurring NA polysaccharde. The term "modified size identification" refer to synthesis of hyaluronic acid molecules of a size distribution not normally found with the native enzyme; the engineered size could be much smaller or fuzer than normal.

[0114] Various hyaluronic acid products of differing size have application in the areas of drug delivery and the generation of an enzyme of altered structure can be combined with a hyaluronic acid of differing size. Applications in angiogenesis and wound healing are potentially large if hyaluronic acid polymers of about 20 monosaccharides can be made in good quantities. Another particular application for small hyaluronic acid oligosaccherides is in the stabilization of the problem of the problem is the collegue of the problem of the blood and a short biological half life. One present solution to this problem is to couple a small molecule shield that prevents the protein from being cleared from the circulation to rapidly. Very small molecular weight hyaluronic acid is well suited for this role and would be nonimmunogenic and biocompatible. Larger molecular weight hyaluronic acid attached to a drug or protein may be used to target the retouloendothelial cell system which has endocytic receptors for hyaluronic acid.

[0116] One of ordinary skill in the art given this disclosure would appreciate that there are several ways in which the size distribution of the hyaluronic acid polymer made by the hyaluronate synthase could be regulated to give different sizes. First, the kinetic control of product size can be altered by decreasing temperature, decreasing lime of enzyme action and by decreasing the concentration of one or both sugar nucleotide substrates. Decreasing any or all of these variables will give lower amounts and smaller sizes of hyaluronic acid product. The disadvantages of these approaches are that the yield of product will also be decreased and it may be difficult to achieve reproducibility from day to day or batch to batch.

[0116] Secondly, the alteration of the intrinsic ability of the enzyme to synthesize a large hyaluronic acid product. Changes to the protein can be engineered by recombinant DNA technology, including substitution, deletion and addition of specific amino acids (or even the introduction of prosthetic groups through metabolic processing). Such changes that result in an intrinsically slower enzyme could then allow more reproducible control of hyaluronic acid size by Kinetic means. The final hyaluronic acid size distribution is determined by certain characteristics of the enzyme, that the operation of prosthetic properties of the enzyme state of production and the enzyme state of production and the enzyme characteristics of the enzyme, that the enzyme can make. Specific changes in any of these residues can produce a modified HAS that produces an HA product having a modified size distribution. Engineered changes to seHAS, spHAS, pmHAS, or cHAS that decreases the intrinsic size of the hyaluronic acid that the enzyme can make before the hyaluronic acid is released, will provide powerful means to produce hyaluronic acid product of smaller or potentially largers size than the native events.

[0117] Finally, larger molecular weight hyaluronic acid made be degraded with specific hyaluronidases to make lower molecular weight hyaluronic acid. This practice, however, is very difficult to achieve reproducibility and one must metuculously reputly the hyaluronic acid to remove the hyaluronidase and unwanted diseasion products.

[0118] As shown in FIG 4, hyaturonan synthase can be engineered to produce hyaturonic acid polymers of different size, in particular smaller, than the normal wildtype enzyme. The figure shows the distribution of FIA sizes (in millions

of Dattons, a measure of molecular weight) for a series of spHAS enzymes, each of which was engineered by site directed mutagenesis to have a single amino acid change from the native enzyme. Each has a different Cysteine residue replaced with Alanine. The cluster of five curves with open symbols represent the following spHAS proteins wildtype, C124A, C261A, C366A, and C402A. The filled circles represent the poorly expressed C225A protein which is only partially active.

[0119] The filled triangles is the C280A spHAS protein, which is found to synthesize a much smaller range of HA polymers than the normal enzyme or the other variants shown. This reduction to practice shows that it is feasible to engineer the hyaluronate synthase enzyme to synthesize a desired range of HA product sizes. The seHAS, pmHAS, and cvHAS genes encoding hyaluronate synthase can also be manipulated by site directed mutagenesis to produce an enzyme which synthesizes a desired range of HA product sizes.

[0120] Structurally, modified hyplatronic acid is no different conceptually than altering the size distribution of the hyplatronic acid product by changing particular amino acids in the desired HAS or the spHAS. Derivatives of UDP-GloNAc, in which the N-acetyl group is missing UDP-GloN or replaced with another chemically useful first storing substants specificily must rely on a particular subset of aniho acids among the 20 is that are conserved. Specific changes to one or more of these residues creates a functional synthese with interactives specifically with one or more of the substantes than the native enzyme. This altered enzyme could then utilize alternatural or special sugar nucleotides to incorporate sugar derivatives designed to allow different chemistries to be employed for the following purposes. (i) covalently coupling specific drugs, proteins, or toxins to the structurally modified hyplatronic acid for general or targeted drug delivery, radiological procedures. ce. (ii) covalently cross linking the yellowing call of the companies of the companies of the companies of the properties and (iii), covalently finking hyplatronic acid to a surface to create a discompatible film or monolayer.

[0121] Bacteria can also be engineered to produce hyaluronic acid. For instance, we have created strains of E. subtilis containing the spHAS gene, as well as the gene for one of the sugar nucleotide precursors. We chose this bacteria since it is frequently used in the biotech industry for the production of products for human use. These bacteria were intended as first generation prototypes for the generation of a bacterium able to produce hyaluronic acid in larger amounts than presently available using a wild type natural stant. We put in multiple copies of these genes.

[0122] For example, three Bacillus subtilis strains were constructed to contain one or both of the Streptococcus propense genes for hyaluronan synthase (sph/48), and UDP-glucose dehydrogenase, the results of which are shown in Table II-8. Based on a sensitive commercial radiometric assay to detect and quantitate HA, it was determined that the strain with both genes (strain #3) makes and secretes HA into the medium. The parent strain or the strain with the dehydrogenase gene (strain #1) does not make HA. Strain #2, which contains just the spHAS gene alone makes HA, but only 10% of what strain #3 makes. Agarose gel electrophoresis showed that the HA secreted into the medium by strain #3 to every high molecular weight.

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Strain Number	Cells	Medium(*)	Strain with genes	Cell density (A ₆₀₀)
		(µg HA per m	ni of culture)	
1	0	0	hasB	4.8
2	4	35	SpHAS	3.9
3	≈>10	>250	SpHAS + has8	3.2

(*) Most HA is in media but some was cell-associated, HA was determined using the HA Test 50 kit from Pharmacia

These experiments used the streptococcal promoters normally found with these genes to drive protein expression. It is expected that the construction of strains with the spHAS or seHAS reading frame under control of a B. subhils promoter would yield even more superior results. The vector used it's a Gram postive/E Colf shuttle vector that has a medium copy number in B. subtitis and a gene for erythromycin resistance (enabling resistance to B µg/mi in B. subtitis not 175 µg/mi in E. coli). The B. subtitis nots strain used is 1A1 from BGSC, which has a tryptophan requirement but otherwise is widitype, and can sporulate. Cell growth and HA production was in Spizizens Minimal Media plus tryptophan, glucose, trace elements and erthromycin (8 µg/mi). Growth was at 32 degrees Celsius with vigorous agitation until the medium was exhausted (36 hours).

[0123] This demonstrates that these bloengineered cells, which would not normally make hyaluronic acid, became competent to do so when they are transformed with the spHAS gene. The seHAS would also be capable of being, introduced into a non-hyaluronic acid producing bacteria to create a bloengineered bacterial strain capable of producing hyaluronic acid.

[0124] A preferred embodiment of the present invention is a purified composition comprising a polypeptide having

an amino acid sequence in accordance with SEQ ID NO. 2 The term "purified" as used herein, is intended to refer to an HAS protein composition, wherein the HAS protein or appropriately modified HAS protein (e.g. containing § IHIS), tail) is purified to any degree relative to its naturally-obtainable state, i.e., in this case, relative to its purify within a prokaryolic cell extract HAS protein may be isolated from Streptococcus, Pasturella, chlorella virus, patient specimens, recombinant cells, infectediassues, isolated subpopulation of issues that contain high levels of hyaluronate in the viracellular matrix, and the like, as will be known to those of skill in the art, in light of the present disclosure. For instance, the recombinant setHAS or spHAS protein makes up approximately 10% of the total membrane protein of E., co. A purified HAS protein composition therefore also refers to a polypetide having the amino acid sequence of SEQ ID NO.2, free from the environment in which it may naturally occur (FIG. 5).

[0125] Turning to the expression of the seHAS gene whether from genomic DNA, or a cDNA, one may proceed to prepare an expression system for the recombinant preparation of the HAS protein. The engineering of DNA segment (s) for expression in a prokaryotic or eukaryotic system may be performed by techniques generally known to those of skill in recombinant expression.

[0128] HAS may be successfully expressed in eukaryotic expression systems, however, the inventors aver that bacterial expression systems can be used for the preparation of HAS for all purposes. It is believed that bacterial expression will ultimately have advantages over eukaryotic expression in terms of ease ef use, cost of production, and quantity of material obtained thereby.

[0127] The purification of streptococcal hyaluronan synthase (seHAS and spHAS) is shown in Table III and FIG. 8. Fractions from various stages of the purification scheme were analyzed by SDS-PAGE on a 12 5% gel, which was then stained with Coomassie Brillhart Blue R-250. Lanes: molecular weight markers; 1, whole E.coli membranes containing the recombinant seHAS-H6; 2, insoluble fraction after detergent solubilization of membranes; 3, detergent solubilized fraction; 4, flow-through from the Ni-NTA chromatography resin; 5-9, five successive washes of the column (two column volumes each); 10, the eluted pure HA synthase which is a single band.

TABLE III

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Step	Total Protein (ug)	Specific Activity (mmol/ug/ hr.	Total Activity (nmol UDP-GlcA)	Yield (%)	Purification (-fold)
Membranes	3690	1.0	3649	100	1.0
Extract	2128	2.2	4725	129	2.2
Affinity Column	39	13	500	14	13.1

[0128] It is proposed that transformation of host cells with DNA segments encoding HAS will provide a convenient means for obtaining a HAS protein. It is also proposed that cDNA, genomic sequences, and combinations thereof, are suitable for eukaryotic expression, as the host cell will, of course, process the genomic transcripts to yield functional mRNA for translation into protein.

[0129] Another embodiment of the present invention is a method of preparing a protein composition comprising growing a recombinant host cell comprising a vector that encodes a protein which includes an amino acid sequence in accordance with SEQ ID NO.2 or functionally similar with conserved or semi-conserved amino acid changes. The host cell will be grown under conditions permitting nucleic acid expression and protein production followed by recovery of the protein so produced. The production of HAS and ultimately HA, including the host cell, conditions permitting nucleic acid expression, protein production and recovery will be known to those of skill in the art in light of the present disclosure of the selfAS gene, and the selfAS gene

[0130] Preferred hosts for the expression of hyaluronic acid are prokaryoles, such as S. equisimilis, and other suitable members of the Streptococcus species. However, it is also known that HA may be synthesized by heterologous host cells expressing recombinant HA synthase, such as species members of the Becilius. Enterococcus, or even Escherichia genus. A most preferred host for expression of the HA synthase of the present invention is a bacteria transformed with the HAS gape of the present invention, such as Laccoccus species, Bacilius suitifies or E coli.

[0331] It is similarly believed that almost any eukaryotic expression system may be utilized for the expression of HAS e.g., baculoivus-based, glutamine synthase-based, dilydrotolate reductase-based systems, SV-40-based, adenovrus-based, cytomegalovirus-based, years-based, and the like, could be employed. For expression in this manner, one would position the coding sequences adj-acent to and under the control of the promoter. It is understood in the art that to bring a coding sequence under the control of such a promoter, one positions the 5° end of the transcription initiation site of the transcriptional reading frame of the protein between about 1 and about 50 nucleotides "downstream" of (i e., 3° of) the chosen promoter. Also, Saccharyoroce aceversize yeast expression vector systems, such as pYES2 via also produced In-Cohen promoter, also, Saccharyoroce aceversize purplied with UPO-Cloka and UPD-GlokNa, or the produced in recombinant yeast-tuding the pYES2 plasmid, When supplied with UPD-GlokNa and UPD-GlokNa, or the produced in recombinant yeast-tuding the pYES2 plasmid, When supplied with UPD-GlokNa and UPD-GlokNa, or the produced in recombinant yeast-tuding the pYES2 plasmid. When supplied with UPD-GlokNa and UPD-GlokNa, or the produced in recombinant yeast-tuding the pYES2 plasmid. When supplied with UPD-GlokNa and UPD-GlokNa, or the produced in recombinant yeast-tuding the pYES2 plasmid. When supplied with UPD-GlokNa and UPD-GlokNa, or the produced in recombinant yeast-tuding the pYES2 plasmid. When purplied with UPD-GlokNa and UPD-GlokNa, or the produced in recombinant yeast-tuding the pYES2 plasmid. When purplied with UPD-GlokNa and UPD-GlokNa, or the produced in recombinant yeast-tuding the pYES2 plasmid. When per produced in recombinant yeast-tuding the pYES2 plasmid. When purplied with UPD-GlokNa and UPD-GlokNa, or the produced in recombinant yeast-tuding the pYES2 plasmid. When purplied with UPD-GlokNa and UPD-GlokNa and UPD-GlokNa produced in recombinant yeast-tuding the pYES2 plasmid. When pro

makes high molecular weight HA.

[0132] Where eukaryotic expression is contemplated, one will also typically desire to incorporate into the inscriptional unit which includes the HAS gene or DNA, an appropriate polyadenylation site (e.g., 5-ATAAA-3) from sever not contained within the original cloned segment. Typically, the poly A addition set is placed about 30 to 2000 nucleotides "downstream" of the termination site of the protein at a opsition prior for transcription termination.

[0133] It is contemplated that virtually any of the commonly employed host cells can be used in connection with the expression of HAS in accordance herewith. Examples of preferred cell lines for expressing HAS cDNA of the present invention include cell lines typically employed for eukaryotic expression such as 239, AtT-20. HepG2, VERO, HeLa, CHO, WI 38, BHK, COS-7, RIN and MDCK cell lines. This will generally include the steps of providing a recombinant host bearing the recombinant DNA segment encoding the HAS enzyme and capable of expressing the enzyme, culturing the recombinant host in media under conditions that will allow for transcription of the cloned HAS gene or cDNA and appropriate for the production of the hyaluronic acid; and separating and purifying the HAS enzyme or the secreted hyaluronic acid from the recombinant host.

[0134] Generally, the conditions appropriate for expression of the cloned HAS gene or cDNA will depend upon the promoter, the vector, and the host system that is employed. For example, where one employs the lac promoter, one will desire to induce transcription through the inclusion of a meteral that will simulate lac transcription, such as isopropytithogalactoside. For example, the cloned selfAS gene of the present invention is expressed as a HIS, containing protein in E. coil as shown in FIG.5. Where other promoters are employed, different materials may be needed to induce or otherwise up-requisite transcription.

[0135] FIG. 5 depicts the overexpression of recombinant seHAS and spHAS in E. coli. Membrane proteins (5mg per lane) were fractionated by SDS-PAGE using a 10% (w/v) get under reducing conditions. The get was stained with Coomassie blue R-250, photographed, scanned, and quantitated using a molecular dynamics personal densirometer (model PDSI P60). The position of HA synthase is marked by the arrow. Lane A is native spHAS (Group A); Lane C is native seHAS; Lane E is recombinant seHAS; Lane P is recombinant spHAS; Lane V is vector alone. Standards used were Bio-rad low Mr and shown in KDs.

[0136] In addition to obtaining expression of the synthase, one will preferably desire to provide an environment that is conducive to HA synthesis by including appropriate genes encoding enzymes needed for the biosynthesis of sugar nucleotide precursors, or by using growth media containing substrates for the precursor-supplying enzymes, such as N-acet/foliucosamine or glucosamine (GicNN-h) and discose (Gic).

30 [0137] One may further desire to incorporate the gene in a host which is defective in the enzyme hybalizonidase, so that the product synthesized by the enzyme will not be degrated in the medium. Furthermore, a host would be chosen to optimize production of HA. For example, a suitable host would be one that produced large quantities of the sugar nucleotide precursors to support the HAS enzyme and allow it to produce large quantities of HA. Such a host may be found naturally or may be made by a variety of techniques including mutageness or recombinant DNA technology.
50 The genes for the sugar nucleotide synthesizing enzymes, particularly the UDP-Glic dehydrogeniae required to produce UDP-Glic, oculd also be isolated and incorporated in a vector along with the HAS gene or CDNA. A preferred embodiment of the present invention is a host containing these ancillarly recombinant gene or cDNAs and the amplification of these ones products thereby allowing for increased production of HA.

[0138] The means employed for culturing of the host cell is not believed to be particularly crucial. For useful datalis, one may wish to refer to the disclosure of U.S. Fat Nos. 45,17,254, 480,159.47,47,48,990, or 47,904,414, all incorporated herein by reference. Where a prokaryotic host is employed, such as S. equisimilis, one may desire to employ a fermentation of the bacteria under anierobic conditions in CO_enriched broth growth media. This allows for a greater production of HA than under aerobic conditions. Another consideration is that Streptococot cellag grown asserboilty do not produce pyropenic exotoxins. Appropriate growth conditions can be customized for other prokaryotic hosts, as will be known to those of skill in the art, inition of the oresent disclosure.

[0139] Once the appropriate host has been constructed, and cultured under conditions appropriate for the production of HA, one will desire to separate the HA so produced. Typically, the HA will be secreted or otherwise shed by the recombinant organism into the surrounding media, allowing the ready isolation of HA from the media by known techniques. For example, HA can be separated from the cells and debris by filtering and in combination with separation from the media by precipitation by alcoholis such as ethanol. Other precipitation agents include organic solvents such as acetine or quaternary organic ammonium salts such as every by prindium orbinder (CPC).

[0140] A preferred technique for isolation of HA is described in U.S. Pat. No. 4,517,295, and which is incorporated at deriven by reference, in which the organic cathodylic acid, inclinionacetic acid, is added to the bacterial suspension at the end of the fermentation. The Inchloroacetic acid causes the bacterial cells to clump and die and facilitates the ease of separating these cells and associated debris from HA, the desired product. The claimfed supernatant is concentrated and dialyzed to remove low molecular weight contaminants including the organic acid. The aforementioned procedure utilizes filtration through filter cassettes containing 0.22 µm pore size filters. Disfiltration is continued until the conductivity of the solution decreases to approximately 0.5 mega-others.

[0141] The concentrated HA is precipitated by adding an excess of reagent grade ethanol or other organic solvent and the precipitated HA is then dired by washing with ethanol and vacuum dired, lyophilized to remove alcohol. The HA can then be redissolved in a borate buffer, pH 8, and precipitated with CPC or certain other organic ammonium salts such as CETAB, a mixed trimethyl ammonium bornide solution at 4 degree(s) Celsius. The precipitated HA is recovered by correse filtration, resuspended in In MACI, dislittered and concentrated as further described in the above referenced patent. The resultant HA is filter sterilized and ready to be converted to an appropriate salt, dry powder or sterile solution, clepending on the desired end use.

A. Typical Genetic Engineering Methods Which May Be Employed

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[0142] If cells without formidable cell membrane barriers are used as host cells, transfection is carried out by the calcium phosphate precipitation method, well known to those of skill in the art. However, other methods may also be used for introducing DNA into cells such as by nuclear injection, cathonic lipids, electroporation, protoplast fusion or by the Biolistictim) Bioparticle delivery system developed by DuPont (1989). The advantage of using the DuPont system is a high transfermation efficiency. If prokaryotic cells or cells withic nothal substantial cell vell constructions are used, the preferred method of transfection is calcium treatment using calcium chloride to induce competence or electropo-

[0143] Construction of suitable vectors containing the desired coding and control sequences employ standard ligation techniques, isolated plasmids or DNA fragments are cleaved, failored, and religated in the form desired to construct the plasmids required. Cleavage is performed by treating with restriction enzyme (or enzymes) in suitable buffer. In general, about 1 µg plasmid or DNA fragments are used with about 1 unit of enzyme in about 20 µl of buffer solution Appropriate buffers and substrate amounts for particular restriction enzymes are specified by the manufacturer, incubation times of about 1 hour at 37° C are workable

[0144] After incubations, protein is removed by extraction with phenol and chloroform, and the nucleic acid is recovered from the aqueous fraction by precipitation with ethanol. If blunt ends are required, the preparation is treated for 15 minutes at 15° C with 10 units of Polymerase (Klenowy, phenol-chloroform extracted, and ethanol precipitated For ligation approximately equimolar amounts of the desired components, suitably end tailored to provide correct matching are treated with about 10 units 17 DNA ligase per 0.5 µg DNA. When cleaved vectors are used as components, it may be useful to prevent religation of the cleaved vector by pretreatment with bacterial skikatine phosphatases.

[0145] For analysis to confirm functional sequences in plasmids constructed, the first step was to amplify the plasmid DNA by cloning into specifically competent E. coil SURTE cells (Stratagene) by doing bransformation at 30-32 C. Second, the recombinant plasmid is used to transform E. coil KS strain Bil337-41, which can produce the UDP-GlotA precursor, and successful transformants selected by antibiotic resistance as appropriate. Plasmids from the library of transformants are then screened for bacterial colonies that exhibit HA production. These colonies are picked, amplified and and the plasmids purified and analyzed by restriction mapping. The plasmids showing indications of a functional HAS gene are then further characterized by any number of sequence analysis techniques which are known by those of ordinary skill in the principle.

B. Source and Host Cell Cultures and Ventors

[0146] In general, prokaryotes were used for the initial cloring of DNA sequences and construction of the vectors useful in the invention. It is believed that a suitable source may be Gram-positive cells, particularly those derived from the Group C Streptococcal strains. Bacteria with a single membrane, but a thick cell wall such as Stephylococci and Streptococci are Gram-positive. Gram-negative bacteria such as E. coli contain two discrete membranes rather than one surrounding the cell. Gram-negative organisms tend to have thinner cell walls. The single membrane of the Gram-positive organisms is a nallogous to the inner plasma membrane of Gram-negative bacteria. The preferred host cells are Streptococcus strains that are mutated to become hysburnoidase negative or otherwise inhibited (EP14016). EP266678, EP244757). Streptococcus strains that have been particularly useful include S equisimits and S zocepidenius.

[0147] Prokaryotas may also be used for expression. For the expression of HA synthase in a form most likely to accommodate high molecular weight HA synthesis, one may desire to employ Streptococcus species such as S. equisimilis or S. zooppidemicus. The aforementioned strains, as well as E. coil W3110 (F., lambda., prototrophic, ATCC No. 273325), bacillis usch as Bacillus subtilis or other enterobacteriaceae such as Serratia mercescens, could be utilized to generate a "super" HAS containing host.

[0148] In general, plasmid vectors containing origins of replication and control sequences which are derived in species compatible with the host cell are used in connection with these hosts. The vector ordinarily carries an origin of replication, as well as marking sequences which are capable of providing phenotypic selection in transformed cells For example. E collis broadful transformed cells are considered to the control of the contro

genes for ampicillin and tetracycline resistance and thus provides easy means for identifying transformed cells. A pBR plasmad or a pUC plasmid, or other microbial plasmid or phage must also contain, no be modified to contain, promoters which can be used by the microbial organism for expression of its own proteins.

[0149] Those promoters most commonly used in recombinant DNA construction include the IacZ promoter, Iac promoter, the T7 bacteriophage promoter, and tryptophan (trp) promoter system. While those are the most commonly used, other microbial promoters have been discovered and utilized, and details concerning their nucleotide sequences have been published, enabling a skilled worker to ligate them functionally with plasmid vectors. Also for use with the present inventor one may utilize integration vectors.

[0150] In addition to prokaryotes, eukaryotic microbes, such as yeast cultures may also be used. Saccharomyces cerevicies or common baker's yeastis the most commonly used among eukaryotic microorganisms, although a number of other strains are commonly available. For expression in Saccharomyces, the plasmid YRp7, for example, is commonly used. This plasmid already contains the frp1 gene which provides a selection marker for a mutant strain of yeast lacking the ability to grow without tryptophan. For example, IRC ON, 4dQ76 or F2P4-1. The presence of the frp1 gene on as a characteristic of the yeast host cell genome then provides an effective environment for detecting transformation by growth in the absence of thyptophan. Suitable promoting sequences in yeast vectors include the promoters for the galactose utilization genes, the 3-phosphoglycerate kinase or other glycolytic enzymes, such as enclase, glyceratide-hyde-3-phosphate dehydrogenase, hexokinase, pyruvate decarboxylase, phosphofutcokinase, glucose-6-phosphate isomerase, 3-phosphoglycora isomerase, and propoglucose isomerate mulsae, pyruvate kinase, time superhosphate isomerase, phosphoglucose isomerase, and propoglucose isomerates.

[0151] In constructing suitable expression plasmids, the termination sequences associated with these genes are also ligated into the expression vector 3' of the sequence desired to be expressed to provide polyadenylation to the mRNA and termination. Other promoters, which have the additional advantage of transcription controlled by growth conditions are the promoter region for alcohol dehydrogenase 2, cytochrome C, said-phosphatase, degradative engrees associated with introgen metabolism, and the aforementioned glyceraldehyde-3-phosphate dehydrogenase, and enzymes responsible for maitose and galactose utilization. Any plasmid vector containing a yeast-compatible promoter, ording of refiliation and termination sequences is suitable.

and glucokinase.

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[0152] In addition to microorganisms, cultures of cells derived from multicellular organisms may also be used as hosts. In principle, any such cell culture is workable, whether from vertebrate or invertebrate culture. However, interest has been greatest in vertebrate cells, and propagation of vertebrate cells in culture has become a routine procedure in recent years. Examples of such useful host cell lines are VERO and HeLa cells, Chinese hamster overy (CHO) cell lines, and Vi38, BHK. COS, and MDCK cell lines.

[0153] For use in mammalian cells, the control functions on the expression vectors are often provided by, viral material. For example, commonly used promoters are derived from polyoma, Adenovirus 2, bovine papilloma virus and most frequently Simian Virus 40 (SV40). The early and late promoters of SV40 virus are particularly useful because both are obtained easily from the virus as a fragment which also contains the SV40 viral origin of replication. Smaller or larger SV40 fragments may also be used, provided there is included the approximately 250 bp sequence extending from the Hild III site toward the Boll visit to loaded in the viral origin of replication.

[0164] Further, it is also possible, and often desirable, to utilize promoter or control sequences normally associated with the desired gene sequence, provided such control sequences are compatible with the host cell systems. An origin of replication may be provided either by construction of the vector to include an exogenous origin, such as may be derived from SV40 or other viral (e.g., Polyoma, Adeno, BPV) source, or may be provided by the host cell chromosome, the precipication mechanism. (If the vector is integrated into the host cell chromosome, the latter mechanism is often sufficient.

C. Isolation of a bona fide HA synthase gene from a highly encapsulated strain of Group C Streptococcus equisimilis.

[0155] The encoded protein, designated seHAS, is 417 amino acids (calculated molecular weight of 47,778 and pl of 9.1) and is the smallest member of the HAS family identified thus far (FIG. 2), seHAS also migrates anomalously fast in SIS-PAGE (M.-42 Vba) (FIGS. 5 and 8).

[0158] FIG. 8 is a graphical representation of a Western Blot analysis of recombinant setAS using specific antibodies Group C (C. Itan 1) or Group A (A.) tan 4) Streptococcal membranes and E. coli membranes (by mglane) containing recombinant setAS (E, lanes 2, 7, and 9) or sptAS (P; lanes 3, 6, 8, and 10) were fractionated by reducing SDS-PAGE and electricitansferred to nitrocellulose. Strips of nitrocellulose were probed and developed as described in the application using purified [36] fractions raised to the following regions of sptAS* central domain peptide E¹¹⁴, T¹⁰⁷ (and 10) and the set of the

[0157] The seHAS and spHAS protein (previously identified in U.S. Serial No. 08/899,940) encoding sequences are 72% identical. The deduced protein sequence of set-HAS was confirmed by reactivity with a synthetic peptide antibody (FIG. 6). Recombinant seHAS expressed in E. coli was recovered in membranes as a major protein (FIG. 5) and

synthesized vary large molecular weight HA in the presence of UDP-GlcNAc and UDP-GlcNa in vitro (FIG. 9). [0158] FIG. 9 shows kinetic analysis of the HA size distributions produced by selfAS and spHAS. E. coff remotranes containing equal amounts of selfAS or spHAS protein were incubated at 37° C with 1.35 mM UDP-[16] GlcA (1.3 x 10³ dpm/nmol) and 3.0 mM UDP-GlcNAc as described in the application. These substrate concentrations are greater than 15 times the respective Km valves. Samples taken at 05, 10, and 60 min were treated with SDS and chromotygraphed over Sephacryl S400 HR. The HA profiles in the fractionation range of the column (fractions 12-24) are normalized to the percent of total HA in each fraction. The values above the arrows in the top panel are the fMWs (in millions) of HA determined directly in a separate experiment using a Dawn multiangle laser light scattering instrument (Wyat. Technology Corp.). The size distributions of HA synthesized by seHAS (P, IL, and aspHAS (O, IL_) at 0.5 min (...), at 1.0 min (...), at 10° strone as indicated. Analysis showed that set HAS and spHAS are essentially identical in the size distribution of HA chains they synthesize (FIG. 9). SeHAS is twice as fast as spHAS in its ability to make HA.

C.1 Bacterial strains and vectors

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[0159] The mucoid group C strain D181; (Streptococcus equisimilis) was obtained from the Rockfeller University Collection. The E. coll host strains Sure and XL1-Blue MRF were from Stratagene and strain Top10 F' was from Invitrogen. Unless otherwise noted, Streptococci were grown in THY and E. coll strains were grown in LB medium. pKK-223 Expression vector was from Pharmacia, PCR 2.1 cloning vector was from Invitrogen, and predigested ½. Zap Express TM Bar HIJCAP Vector was from Stratagene.

C.2 Recombinant DNA and Cloning

[0160] High molecular mass Genomic DNA from *Streptococcus equisimitis* isolated by the method of Caparon and Scott (as known by those with ordinary skill in the art) was partially digested with SauSA1 to an average size of 2-12 kb. The digested DNA was precipitated with ethanol, washed and ligated to the *Bam HI/CIAP)*. Zap Express vector Ligated DNA was packaged into phage with a Packagene^{1M} extract obtained from Promega. The titer of the packaged phage library was checked using XL1-filtus MFFT E. coil as a host.

30 C.3 Degenerate PCR Amplification

[0161] Degenerate oligonucleotides were designed based upon conserved sequences among spHAS (Straptococus pyageness). DG42 (Xenopus leavis HAS; 13) and node a Rhizobium melilotin nodulation factor, 20) and were used for PCR amplification with D181 genomic DNA as a template. Amplification conditions were 34 cycles as til 54°C for 1 min, A4°C for 1 min, 72°C for 1.5 min followed by a final extension at 72°C for 10 min. Oligonucleotide HADRF1.5°GAY MOA YRT YTA CX AAT TAY GCT ATH GAY TIR G9-3' (SEQ ID NO 20), sense strand) corresponds to the sequence D²⁴⁸RCLTHYADLC (SEQ ID NO 9; spHAS). Oligonucleotide HACRF1.5°GACG WGT WCC CCA NTC XGY ATT TIT AD XGT RCA* (SEQ ID NO 10; an intense strand) corresponds to the region C6*4TIKNTEWDTRT (SEQ ID NO 10; spHAS). The degeneracy of bases at some positions are represented by nomenclature adopted by the IUPAC in its codes for despenerace bases lighted in Table IV.

TABLE IV IUPAC Codes - Degenerate Bases

The International Union for Fure and Applied Chemistry (IUPAC) has established a standard single-letter designation for degenerate bases. These are:

C+G+T В D A+G+T H A+C+T ĸ T+G A+C M. == N A+C+G+T R A+G S ,,, G+C W T+A U

V = A+C+G
X = a minor bases (specified elsewhere)
Y = C+T

C.4 Library Screening

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[D162] Two molecular probes were used to screen the library, the cloned 459 bp PCR product and oligonucleotide D181.5 (5-CCTTGATAGGTGACACGTGTGACG-3 (SECI ID No. 15); derived from the 1042 by sequence). The 453 bp PCR product was radiolabeled using the Prime-It 11 random primer labeling Kit (Stratagene) according to the manufacturers instructions. Oligonucleotides were labeled by Kinace-It Kinasing Kit (Stratagene) using [P2F]ATP. Radiolabeled products were separated from noniabeled material on Nucl*rap Push columns (Stratagene) and P161 genomic digest on Southern blots. To screen the 2-phage library, XLB-UE MRF was used as a hot (3000 plaques/plate) on Ntinocellulose membranes containing adsorted phage, were prehyribridate at 60°C according to screen the 2-phage library. XLB-UE MRF was used as a hot (3000 plaques/plate) on Ntinocellulose membranes containing adsorted phage, were prehyribridate at 60°C according to instructions.

[0163] The membranes were then washed with 2x SSC buffer and 0.1% (w/v) SDS at room temperature for 15 min. at 80°C with 0.1x SSC buffer and 0.1% SDS (w/v) for 30 min, dired and then exposed to Bio-Max MS film overnight at -70°C. Positive plaques were replated and rescreened twice. Pure positive phages were saved in SM buffer with chloroform. PCR on these phages with vector primers revealed 3 different insert sizes.

(0164) PCR with a combination of vector primers and primers from different regions of the cloned 1042 bp sequence revealed that only one of the three different phages had the complete HAS gene. The insert size in this phage was 6.5 kb. Attempts to subclone the insert into plasmid form by autoexcision from the selected phage library clone failed. Therefore, a PCR strategy was applied again on the pure positive phage DNA to obtain the 5" and 3" end of the OR [Olioonucleotide primers DFAI 3.6"-GCCCTGTGTACGACTICA-3" (SEQ ID NO.16.1) and 7.3" (vector primer).

plified a 3kb product and oligonucleotides D181.5 and T7 (vector primer) amplified a 2.5 kb product. The 5' and 3'-end sequences of the ORF were obtained by sequencing these two above products. Analysis of all PCR product sequences allowed us to reconstruct the ORF of the 1284 bp seHAS gene.

C.5 Expression cloning of the seHAS

[0163] Primers were designed at the start and stop codon regions of seHAS to contain an EcoR1 restriction site in the sense oligonuclectide (5°-AGGATCCGAATTCATGAGAACATC-3′-(EGC ID NO.171)) and a PSf1 site in the antisense oligonuclectide (5°-AGAATTCTGCAGTTATAATAATTTTTACGTGT-3′-(EGC ID NO.18)). These primers amplified a 1.2 kb PCR product from D181 genomic DNA as well as from pure hybridization-positive phage. The 12 kb product was purified by agarose gel electrophoresis, digested with Psf1 and EcoR7 and cloned directionally into Psf1-and EcoR7-digested pKK223 vector. The ligated vector was transformed into E. coli SURE colls that were there grown at 30°C. This step was practically important since other host cells or higher temperatures resulted in delations of the cloned insert. Colonies were isolated and their pDNA purified. Out of six colonies (named a, b.c.d.e, and fi. five had the correct size insert, while one had no insert.

C.6 HA Synthase Activity

[0168] HA synthase activity was assayed in membranes prepared from the 5 above clones. Fresh log phase cells were harvested at 3000g, washed at 4°C with PBS and membranes were isolated by a modification of a protocolast method as known by those of ordinary skill in the art. Membrane preparations from Streptococcus goujesimilis were also obtained by modification of a different protoplast procedure. Membranes were incubated at 3°C in 50 mM sodium and potassium phosphate, pH 7°C with 20 mM MgCh, 1 mM OTE, 120 jM UDP-GIGA and 300 jM UDP-GIGNAc. Incorporation of sugar was monitored by using UDP-[1*C)GIGA (318 mCimmol; ICN) and/or UDP-GIGNAc. Incorporation of sugar was monitored by using UDP-[1*C)GIGA c 20.3 Climmol KEN]. Reactions were terminated by addition of 50 Sto ta final concentration of a CV (w/v). Product HA was separated from precursors by descending paper chromatography and measured by determining nonomorated radioactivity at the origin.

C.7 Gel Filtration Analysis

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[0167] Radiolabeled HA produced in vitro by membranes containing recombinant seHAS or spHAS was analyzed by chromatography on a column (0.9 x. 40 cm) of Sephacryl \$500 HR (Pharmacia Biotech Inc.). Samplee (0.4 ml in 200 mlM NaCl, 5 ml Tis-HCl, pH 8.0, plus 0.5% \$650) were elucted with 200 mlM NaCl, 5 ml Tis-HCl, and pH 8.0 and 0.5 ml fractions were assessed for ¹⁴C and/or ³H radioactivity. Authenticity of the HA polysaccharide was assessed by treatment of a separate identical sample with the HA-specific hyaluronate lyase of *Streptomyces hyalurolyticus* (EC 4.2.2.1) at 37°C for 3 hrs. The dispest was then subjected to get filtration.

C.8 SDS-PAGE and Western Blotting

[0188] SDS-PAGE was performed according to the Laemmili method. Electrotransfers to nitrocellulose were performed within standard blotting buffer with 20% methanol using a Bio-Rad mini Transblot device. The blots were blocked with 2% BSA in TBS. Protein A/G alkaline phosphatase conjugate (Pierce) and p-nitroblue tetrazolium/5-bromo-4-chloro-3 inddly phosphate p-clubuldine salt were used for detection.

45 C.9 DNA Sequence and Analysis

[0168] Plasmids were sequenced on both strands using fluorescent labeled vector primers. Sequencing reactions were performed using a Thermosequenase™ kit of fluorescent labeled primers (with 7-deazaG). Samples were electrophoresed on a Pharmacia ALF Express DNA Sequencer and data were analyzed by the ALF Manager Software v3 02. Internal regions of inserts were sequenced with internal primers using the ABI Prism 377 (Software version 2.11). Ambiguous regions ever sequenced manually using Sequenase™ 7-deaza - DNA polymerase, 7-deaza GTP master mix (USB) and [tr.²⁵S] dATP (Amersham Life Sciences). The sequences obtained were compiled and analyzed using DNASIS, v2.1 (Hitachi Software Engineering Co., Ltd.). The nucleotide and amino acid sequences were compared with other sequences in the Genbank and other databases.

C.10 Identification of seHAS

[0170] Identification of seHAS was accomplished by utilizing a PCR approach with oligonucleotide primers based

on several regions of high identity among spHAS, DG42 (now known to be a developmentally regulated X. Isevis HAS and designated xt HAS) and NodC (a Rhizobium B-G10Ako transferase). The xtHAS and NodC proteins are, respectively. 50% and -10% identical to spHAS. This strategy yielded a 459 bp PCR product whose sequence was 66.4% identical to spHAS, indicating that a Group C hornologue (seHAS) of the Group A (spHAS) HAS ynthase gene had been identified. The complete coding region of the gene was then reconstructed using a similar PCR-based strategy. A final set of PCR primers was then used to amplify the complete ORF from genomic DNA. When this 1.2 kb PCR fragment was incorporated into the expression vector pKK223 and transformed into E. coli SURE cells, HA synthetic activity was demonstrated in isolated membranes from 5 of the Solonies tested.

[0171] The ORF of the reconstructed gene encodes a novel predicted protein of 417 ammo acids that was not in the database and it is two armio acids shorter than spHAS. The two bacterial proteins are 72% identical and the notice acid sequences are 70% identical. The predicted molecular weight of the seHAS protein is 47.778 and the predicted isoelectric point is at pH 9.1. Three recently identified mammalian HASs (muHAS2, muHAS2, muHAS3, RIG. 2) are similar to the bacterial proteins. The overall identity between the two groups is -28-31%, and in addition many ammo acids in seHAS are highly conserved with those of the eutraryoic HAS6 (e.g. KIR or DIE substitutions). ASBR, the PBCY1-HAS are 28-33 percent identical the mammalian HASs, and is predicted to have a similar topology in the lipid membrane. Within mammalian species the same family members are almost completely identical (e.g. muHAS1 and buHAS2 are 98% identical, muHAS2 and soft-inicial). However, and as shown in FiG. 3. even within the same species the different HAS family members are more divergent (e.g. muHAS1 and muHAS2 are for this dentical).

[0172] FIG. 10 shows hydropathy plots for seHAS and predicted membrane topology. The hydrophilicity plot for the Streptococcal Group C HAS was generated by the method of Kyte and Doolittle (J. Mol. Biol. 157, 105, 1982) using DNAsis. The protein is predicted to be an intereal membrane protein.

[0173] FIG. 11 shows a model for the topologic organization of seHAS in the membrane. The proposed topology for the protein conforms to the charge-in rule and puts the large central domain inside. This domain is likely to contain most of the substate binding and catalytic functions of the enzyme. Cy5²⁷⁶ in seHAS, which is conserved in all HAS family members, as well as the other three cysteines are shown in the central domain. Cys²⁸¹ is a critical residue whose attention can dramatically later the size distribution of HA product synthesized by the enzyme.

[0174] The overall membrane topology predicted for seHAS is identical to that for spHAS and the eukaryotic HASs reported thus far. The protein has two putative transmembrane domains at the amino terminus and 2-3 membrane associated or transmembrane domains at the carboxyl end. The hydropathy plots for the two Streptococcal enzymes are virtually identical and illustrate the difficulty in predicting the topology of the extremely hydrophobic region of -90 residues at 13°13.74°6° in seMAS (X¹³1-X¹⁴0° in seMAS (X¹³1-X¹⁴0° in seMAS (X¹³1-X¹⁴0°) in seMA

[0175] seHAS was efficiently expressed in *E. coli* cells. Roughly 10% of the total membrane protein was seHAS as assessed by staining of SDS-PAGE gels (FIG. 5). The prominent seHAS band at 42 kD is quantitatively missing in the vector-only control lane. This unusually high level of expression for a membrane protein is also found for spHAS, using the same vector in SURE cells. About 9% of the membrane protein is spHAS in *E. coli* SURE cells. About 9% of the membrane protein is spHAS in *E. coli* SURE cells. In contrast, the amount of seHAS in Group C membranes is not more than 19% of the total membrane protein. The spHAS in Group A membranes is barely detectable. The recombinant seHAS expressed in *E. coli* SURE cells does not synthesize HA in vivo since these cells lack UDP-GlcA, one of the required substrates. Membranes, however containing the recombinant seHAS protein synthesize HA when provided with the substrates SUP-GlcAN and UDP-GlcA (FIG. 12).

[0178] FIG. 12 shows the synthesis of authentic HA by recombinant seHAS. E. col/membranes (69 µg) prepare from cells containing recombinant seHAS or vector alone were incubated at 37°C for 1 hour with 700 µM UDP-[1*H] GloNAG (278 x 10* dpm/hmbl. (1) may and 300 µM UDP-[1*C]GloA (3.80 x 10* dpm/hmbl; C.) in a final volume of 200 µl as described herein. The enzyme reaction was stopped by addition of EDTA to a final concentration of 25 mM. Half the reaction mix was treated with Streptomyces hysteroridaes e175°C for 3 hours. SSB (28, wh) was added to hysteroridaes e180 (20, 10* and untreated (1.80* samples, which were heated at 90°C for 1 min. The samples were diluted to 500 µl with column buffer (6 mM Tris, 0.2 M Nacl, pH 8.0), clarified by centrifigation and 200 µl was injected or 3 Sephacryl S-500 HR column. Fractions (1 ml) were collected and radioactivity was determined BD is the peak elution position position of blue destrain (2× 10° DA, Pharmacia), V, marks the excluded volume and V; the included volume. The ratio of (1*4c) GloA: (2*H) GloNAc incorporated into the total amount of HA fractionated on the column is 1.4, which is identical to the ratio of specific activities of the was usustrates. Therefore, the molar ratios of the sugar incorporated into product is 1.1 as predicted for authentic HA. Membranes from cells transformed with vector alone did not synthesize

[0177] Using 120 µM UDP-GicA and 300 µM UDP-GicNAc. HA synthesis was linear with membrane protein (et S.O. 2 µg) and for at least 1 hour. Also, membranes prepared from nontransformed cells or cells transformed with vector alone have no detectable HAS activity. HA synthesis is negligible if Mg⁻² is chelated with EDTA sS% of control) or if either of the two substates are omitted (-2% of control). Recombinant setHAS also showed the expected specificity or sugar nucleotide substrates, being unable to copplymenze either UDP-GalA. UDP-GalVAc with either

of the two normal substrates (Table II).

[0178]. Based on gel filtration analysis, the average mass of the HA synthesized by seHAS in isolated membranes is 5-10x10°D. The product of the recombinant seHAS is judged to be authentic HA based on the equimolar incorporation of both sugars and its sensitivity to degradation by the specific Streptomyces hyaluronidase (FIG. 12). Although the conditions for total HA synthesis were not optimal (since-90% of one substrate was incorporated into product), the enzyme produced a broad distribution of HA chain lengths. The peak fraction corresponds to an HA mass of 7.5x10°D as which is a polymer containing approximately 36,000 monomeric sugars. The distribution of HA sizes resolved on this column ranged from 2-20x10°D as

[0179] The deduced protein sequence of selfAS was confirmed by the ability of antibodies to the spHAS protein to cross-react with the Group C protein (FIG. 8). Polyclonal antibodies to the whole spHAS protein or to just the central domain of spHAS also reacted with the selfAS protein. Antipeptide antibody to the C-terminus of spHAS did not cross-react with this somewhat divergent region in the selfAS protein. However, antipeptide antibody directed against the spHAS sequence E14** T-16** recognized the same predicted sequence in selfAS. The antipeptide antibody also reacts with the native selfAS and spHAS proteins in Streptococcal membranes and confirms that the native and recombinant enzymes from both species are of identical size. Like the spHAS protein, selfAS ingrates anomalously fast on SDS-PAGE. Although the calculated mass is 47.778 b.a. the Ny SDS-PAGE is consistently "42 kDs.

[0180] Because of the sequence identity within their central domain regions and the overall identical structure predicted for the two bacterial enzymes, the peptide-specific antibody against the region E^{1,4,7} T.⁶¹ can be used to normalize for HAS protein expression in membranes prepared from cells transformed with genes for the two different enzymes. Using this approach, membranes with essentially identical amounts of recombinant spHAS or seHAS were compared with respect to the intital rate of HA synthesis and the distribution of HA product size.

[0181] As shown for spHAS, the synthesis of HA Chains by seHAS is processive. The enzymes appear to stay associated with a growing HA chain until it is released as a final product. Therefore, it is possible to compare the rates of HA elongation by seHAS and spHAS by monitoring the size distribution of HA chains produced at early times, during the first round of HA chain synthesis. Based on gel filterition analysis of HA product sizes at various times, we estimated that the average rate elongation by seHAS is about 9.000 monosocchardes/finulted 13.7° C/FIG. 9). In five minister, the enzymes can polymerize an HA chain of 5-10x10°Da. During a 90 min incubation, therefore, each enzyme molecule could potentially initiate, complete and release on the order of 5-5 such larger HA molecules. At early times (a₂ -11 min), reflecting elongation of the first HA chains, the size distribution of HA produced by seHAS was shifted to larger species compared to spHAS. 9, 90 min the two distributions of HA product sizes are indistinguishable.

[0182] The cloned seHAS represents the authentic Group C HA synthase. Previously reported or disclosed "Group C" proteins are, therefore, not the true Group C HAS. The seHAS protein is homologous to nine of the currently known HA synthases from bacteria, vertebrates, and a virus that now comprise this rapidly growing HA synthase family. This homology is shown particularly in FIG. 2. In mammats three genes, designated HAS 1. HAS 2 and HAS 3, have been identified and mapped to three different chromosomes in both human and mouse. In amphibians the only HAS protein identified thus far is the developmentally regulated GG42, which was cloned in 1989 and recently shown to encode the HA synthase activity by analysis of the recombinant protein in yeast membranes. Probably other X. laevus HAS genes will soon be identified.

[0183] A divergent evolution model suggests that a primitive bacterial HAS precursor may have been usurped early during vertebrate development or the bacterial pathogenic strategy of making an HA capsule was developed when a primitive bacteria captured in primordial HAS. Convergent evolution of the bacterial and eukaryotic HAS enzymes, to a common structural solution 'seems unlikely, but may have occurred.

[0184] None of the three mammalian isozymes for HAS have yet been characterized enzymatically with respect to their HA product size. At least, ten identified HAS proteins are predicted to be membrane proteins with a similar topology HA synthesis occurs at the plasma membrane and the HA is either shed into the medium or remains cell associated to form the bacterial capsule or a eutaryotic pericellular roat. The sugar nucleoide substrates in the cytoplasm are utilized to assemble HA chains that are extunded through the membrane to the external space.

[0183] The protein topology in the very hydrophobic carboxyl portion of the HAS protein appears to be critical in understanding how the enzymes extend the growing HA chain as it is simultaneously extuded through the membrane For example, the unprecedented enzymatic activity may require unusual and complex interactions of the protein with the high dislayer. Preliminary results based on analysis of syHAS-aktainer posphatases fusion proteins indicate that the amino and carboxyl termini and the large central domains are all intracellular, as shown in FIGS. 10 and 11. The seHAS protein also contains as large central domain (~63% of the total protein) that appears to contain the two substrate binding attes and the two glycosyltransferase activities needed for HA synthesis. Although current software programs cannot reliably predict the number or nature of membrane-associated domains within the long C-terminal hydrophobic stretch, the proposed topological arrangement agrees with the present evidence and applies as well to the eukarboic enzymes, which are ~40% larger primarily due to extention of the C-terminal end of the protein with 2 additional oredicted transmembrane domains.

[0186] Four of the six Cys residues in spHAS are conserved with seHAs. Only Cys225 in both bacterial enzymes is conserved in all members of the HAS family. Since sulfhydry freactive agents, such as p-mercurobenzoate or NEM, greatly inhibit HAS activity, it is likely that this conserved Cys is necessary or important for enzyme activity, Initial results from site-directed mutagenesis studies, however, indicate that a C225S mutant of spHAS is not inactive, it retains 5-10% of wildfulow activity.

[0187] The recognition of nucleic acid sequences encoding only seHAS, only spHAS, or both seHAS and spHAS using specific oligonucleotides is shown in FIG. 13. Three pairs of sense-antisense oligonucleotides were designed based on the sequence of 10 SEQ NO. 1 and the coding sequence for spHAS. The seHAS based nucleic acid segments (sel-se2 and sesp1-sesp2) are indicated in FIG. 14. These three oligonucleotide pairs were hybridized under typical POR reactions with genomic DNA from either Group C (seHAS) planes 2, 4, and 6) or Group A (spHAS) (lanes 3,5, and 7) streptococi. Lanes 1 and 8 indicate the positions of MW standards in bt (slobases). The POR reactions were performed using Tag DNA polymerase (from Promega) for 25 cycles as follows: 94 degrees Celsius for 1 minute to achieve DNA denaturation, 43 degrees Celsius (42 degrees Celsius for fibe smaller common sesp primers) for 1 minute to allow hybridization, and 72 degrees Celsius for 1.5 minutes for DNA synthesis. The POR reaction mixtures were then separated by electrophoresis on a 1% agrance gel.

[0188] The sal-se2 primer pair was designed to be uniquely specific for the Group C HAS (seHAS). The sp1-sp2 primer pair was designed to be uniquely specific for the Group A HAS (spHAS). The sesp1-sesp2 primer pair was designed to hybridize to both the Group A and Group C HAS nucleic acid sequences. All three primer pairs behaved as expected, showing the appropriate ability to cross-hybridize and support the generation of PCR products that were specific and/or unique.

[0189] The oligonucleotides used for specific PCR or hybridization are shown in FIG. 14. The synthetic oligonucleotides of SEQ ID NOS. 3, 4, 5, and 6 are indicated in the corresponding regions of SEQ ID NO. 1. These regions are in bold face and marked, respectively as primers set, sec_2 sep1, and sesp2. The #1 indicates primers in the sense direction, while the #2 indicates a primer in the antisense direction. Each of the four oligonucleotides will hybridize specifically with the selfAS sequence and the appropriate pairs of sense/antisense primers are suitable for use in the polymerase chain reaction as shown in FIG. 13.

[0190] FIG. 7 shows a gel filtration analysis of hyaturonic acid synthesized by recombinant HAS expressed in year membranes. A DNA fragment encoding the open reading frame of 419 amino acid residuse scoresponding to spHAS (with the original Vel codon switched to Met) was subcloned by standard methods in the pYES2 yeast expression vector (from invitrogen) to produce pYES/HA Membranes from cells with this construct were prepared by a gittation with plass beads. The samples derived from pYES/HA constructs contained substantial HA synthase activity and the "42 KDa" HAS protein was detected by Western analysis using specific antibodies; membranes from cells with vector alone possessed neither activity nor he immunoreactive band (not shown). Membranes (316 up protein) were first included with carrier free UDP-("4C) GIcA (1 uci."4C) amd 900 ub unlabeled UDP-GIcNAo in 50 mM Tris, pH 7, 20 mM MgC12. HMD DTT, and 0.05 M NaCl (450 ul reaction volume) at 30 degrees Celsius for 1.5 minutes. After this pulse-label period normadolabeled UDP-GIcNAo was then added to final concentrations of 900 uM. Samples (100 uL) were taken after the pulse at 1.5 min (dark circle), and 15 (black square), and 45 (black triangle) min after the "chase." The reactions were terminated by the addition of 950s to 2% and heating at 95 degrees Celsius for 1 min. The samples were clarified by centrifugation (10,000 x, 5 min) before injection of half of the sample onto a Sephacryl S-500HR gel filtration column (Pharmacia: 1, 450 cm) equilibrated in 0.2 M NaCl.5 mM Tris, bit 8.

[0191] The column was eluted at 0.5 ml/min and radioactivity in the fractions (1 ml) was quantitated by liquid scintillation counting after adding BioSaf ell cocktail (4.5 ml, Research Products Int1.). The void volume and the totally included volumes were at elution volumes of 14 ml and 35.5 ml, respectively. The peak of blue dextran (average 2x10 6 Da) eluted at 25.27 ml. The recombinant HAS expressed in the eukaryotic yeast cells makes high molecular weight invaluronic acid in vitro.

[0192] Thus it should be apparent that there has been provided in accordance with the present invention a purified nucleic acid segment having a coding region encoding enzymatically active HAS, methods of producing hydronic acid from the setHAS gene, and the use of hyaluronic acid produced from a HAS encoded by the setHAS gene, that fully satisfies the objectives and advantages set forth above. Although the invention has been described in conjunction with specific embodiments thereof, it is evident that many alternatives, modifications, and variations will be apparent to those skilled in the art. Accordingly, it is intended to embrace all such alternatives, modifications, and variations that fall within the spirit and broad scope of the appended claims.

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	1 S 10 15	

	tat	caa	tta	gca	ctc	asa	tta	ttt	gaa	azg	tog	gcg	gaa	atc	tat	gga	egg	aaa	stt	gtt	123
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	Thr	caa Gln	Leu	Leu	Leu	Ser	Asn	Val	Lys	Еў'я 85	Leu	Val	Leu	Ser	Asp 90	Ser	Clu	Lys	Asn	Thr 95	303
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	Trp	gtt Val	Hîs	Ser	7yr 260	Val	A) a	Glu	Leu	Leu 265	Glu	Asp	Asp	Asp	270	Thr	11e	116	GIÀ	275	843
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55	Se:	r Pro	t tte	e egt	ttt Pho	Phe	geg Ala	geg Ala	ggt	aat Asr 34!	1 441	get	Phe	gct	. aaa Lys 350	· cys	tgg	Ct a	aat Aan	322 FA3 949	1083

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					aca Thr 480																1503
25					atg Het 500				Asn												1563
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5	agt Ser	att Ile	ttt Phe	īyr Tat	ecc Pro 720	aat Asn	aca Thr	ita Leu	Хэņ	ggc G1y 725	tta Leu	gtg Val	aaa Lys	aaa Lys	cta Leu 730	aac Asn	aat	att Ile	att	gaa Glu 735	2223
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		atg																			2703
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	Thr	Ser	CAs	Gln	Ser	He	Asp	Ser	Val	Pro		Tyr	Ýsu	Thr	Glu 910		Ile	Trp	Phe	915	
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	Phe	Ala	Leu	Len	Ile 920	Leu	Glu	Lys	Lys	Thr	CTA	His	Val	Phe	930	Lys	Thr	ser	THE	935	
40																					
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Claims

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- An isolated enzymatically-active hyaluronan synthase that is encoded by a nucleic acid sequence that has at least 80% identity with SEQ ID NO: 8.
- The isolated enzymatically-active hyaturonan synthase as claimed in Claim 1, which is encoded by a nucleic acid sequence which has between 80% and 90% identity with SEQ ID NO: 8.
- The isolated enzymatically-active hyaluronan synthase as claimed in Claim 1, which is encoded by a nucleic acid sequence which has between 90% and 99% identity with SEQ ID NO: 8.
 - The isolated enzymatically-active hyaluronan synthase as claimed in Claim 1, which comprises the amino acid sequence of SEQ ID NO; 7.

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- The isolated enzymatically-active hyaluronan synthase as claimed in Claim 1, which consists of the amino acid sequence of SEQ ID NO: 7.
- The isolated enzymatically-active hyaluronan synthase as claimed in any one of Claims 1-5, which is a chlorella virus polypeptide.
 - The isolated enzymatically-active hyaluronan synthase as claimed in any one of Claims 1-6, which is a paramecium bursaria chlorella virus polypeptide.
- An isolated nucleic acid segment encoding the enzymatically-active hyaluronan synthase as claimed in any one
 of Claims 1-7.
 - 9. The nucleic acid segment as claimed in Claim 8, which has the sequence of SEQ ID NO: 8.
- 15 10. A recombinant vector comprising the nucleic acid segment as claimed in Claim 8 or 9, wherein the recombinant vector is a plasmid.
 - 11. A recombinant vector comprising the nucleic acid segment as claimed in Claim 8 or 9, wherein the recombinant vector is a replicating or integration plasmid, cosmid, phage, or virus vector.
 - A recombinant host cell comprising the recombinant vector as claimed in Claim 10 or 11, wherein the host cell produces hyaluronic acid.
- 13. The recombinant host cell as claimed in Claim 12, further comprising at least one precursor gene found in a biosynthetic pathway of hyaluronic acid.
 - The recombinant host cell as claimed in Claim 12 or 13, wherein the host cell is a Bacillus, E. Coll. Lactococcus, Streptococcus or Entergopous host.
- 30 15. The recombinant host cell as claimed in Claim 12 or 13, wherein the host cell is Bacillus subtilis.
 - 16. A method for producing a hyaluronic acid polymer, comprising
- (a) growing the recombinant host cell as claimed in any one of Claims 12-15 in a medium to secrete hyakuronic acid; and (b) recovering the hyakuronic acid.
 - (b) reserving the righterine deta

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- 17. The method as claimed in Claim 16, wherein the step of recovering the hyaluronic acid further comprises the step of extracting the secreted hyaluronic acid from the medium.
- 18. The method as claimed in Claim 17, further comprising the step of purifying the extracted hyaluronic acid.

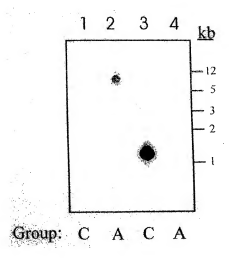


Fig. 1

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hullAS	HHCERFICIL RIIGTTLFGVSLE LGITANHIVG	33
×1HAS	NG-ENTITH VSWITITSNL IAVGGASLIË APAITGNUK HRIERMLIT	49
CYBAS	whialsti wgvsatcifu fgfflagulf selnrkalrk wislrykgwu	87
seRAS	YLFGAKG SLSIGFIL IAYLUVKISL SFF-YKPFKG RAGQY YLFGT-S TYGIYGVIL ITYLVIKLGL SFL-YEPFKG HFHDY YOFIQFDNYY FSFGLYAFL ASHLIIQSLF AFLEHRKHKK SLETPIKL	
	13504KG2F21ZGEFD IMITDAMIZE 265-IVEEKG HMOO	.02
SARgs	TLEGT-STVGIXGVIL ITTLVIKLGL SEL-TEPFKG HPHDY	64
HUHAS	YOFIQTONYY FSFSLYGAFL ASHLIIQSLF AFLEHRRHKK SLETPIKL	B 3
×1HAS	FQVLKHEAIL FSLGLTGLAN LLHLMMQSLF AFLEIRRVNK 5-ELPCSF	96
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CVHAS	HARVIKAIYN DRIKKPEPV ICESDDKEGE	165
	DAY 184114 DAY	
SCHAS		140
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×inas	HMENFKOVER GEDVGTYVWK GRYHTVKKPE ETHRGSCPEV SKPLNEDEGI	196
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SEHAS	HA	163
SAHAS	RA DAY	162
huHAS	CHYTOLYLSH KSICINOKHG GKREVHYTAK RALGRSVD YVOVCDSDEN	215
KIBAS	MANUFACTURE REPORTED CARRIES OF TACABLE AND ARREST AND ARREST OF THE PROPERTY	244
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SAHES	INDUSTRICE KIENDELALV KIR-HTNAM KOLNETART DISKONDERA	212
BAHGE	TYPHALEELL KSPHDETVYA ATG-HLNARH ROTHLLTRLT DIRECHLEGV	211
huHAS	Lekdrikov yplacdpeid avageckimi t-dtilsilv avitiksku Ithalieli ktemptiva atg-hlanni admilitrit diroksku Ithalieli ksembetvia atg-hlanni admilitri diroksku Lopassvým kvileddang gvgdvojik ktosistels svájnakki Loelatvým kvilendnia avgddvali tustietse slejnakki	266
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SPHAS	TRANSETON ILVESTIESI YRREVIIPEL ERYKNITELE LPVSICEDRIC	261
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CVHAS .	THEITHER KAALLANAG REDERINGER LEAVING BACKETALL	361
SCHAS	LTHYATOLG- KTVYQSTARC ITDVHUKHST YLKQQNEYNK KEFFESIISV	311
SPHAS	LTHYAIDLE- RTVYOSTARC DTDVPFOLKS YLKOONEGER SFFRESIISV	310
huHAS	Incilhor xvptepau sosstiner vivotrost buffinist Linkadde ktvostar: itdvidnist ylkonisne Linkadde ktvostar: otdvijolks ilkonisne septistisv Linkasde ktvostar otdvijolks ilkonisne septistis Linkasde ktrtharsk ietesslika blogitasi septimina Linkasde ktrtharsk tetesslika blogitasi septimina	366
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CVHAS	FARMHOLSG INLAFECLYG ITYFFLVIYL FSRLAVEADP RAGIATVIVS	411
SCHAS	KKIKHN9FVA LHTILEVSKF MHLYYSVVDF FVGNVREFDW LRVLAFLVII	361
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huNAS	KKIEMPEVA LETIESVYK MELYSVYOF FYGWARFOM LEVIZELVII KKIEMPEVA LETIESVYK MELYSVYOF FYGWARFOM LEVIZELVII KRIEMPEVA LETIESVYK MELIVATGKE LEPGATGOD LEVIZELVII METRIKKE— LANTESVYS FIFPFFITAT VIRLYAGTI MELYWELLCI OMHRUN— 1MTTESVYS FIFPFFITAT VIRLYAGTI MENYWELLCI	413
×1HA5	DAGGERDA TOMPY COME PIEDFFITH VIRITYAGTI WOUNTITOT	441
Vruug.	**************************************	
	Water Barrier Barrier Barrier Barrier Barrier Barrier	
CVHAS	TTVAJIRGGY ESFARODRA FYFY-LÜTTV YFFCHIĞARI TAVGÜLDICI FYVAJCHILB YHKKIPLE FILLSFÖGUL HEFUDÜLEL YSLĞIRDING FYVAJCHINI YHVERRAS FILLSFÖGUL HEFUDÜLEL YSLĞIRDIF OLVGİRSS- FASCLAGRIY HVERSLÖYÜL YHSLİĞINI FALAİTIKKA OLYGİRSSI YACLIRGEYI HLIMSLÖĞÜN YHTÜLİLİŞIY FALAĞIRKI OLYGİRSSI YACLIRGEYI HLIMSLÖĞÜN YHTÜLİLİŞIY FALAĞIRKI	450
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SPHAS	FIVAECRNUM YMVRHPAS FLLSPLZGIL HLEVLOZEKL YSLCZIENTE	. 4SB
BURAS	OLYGÜIKSS- FASCLAGNIY HYFMSLÜSYL YMSSLLÜKM FAIATINKAG	462
KIHAS	OTHERES - VECKLEGNET HILMSLYSBL YMTGLLESKY FALLTLIKTG	490
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Senas	MCZBZK F	417
SARge	HCL HKK A IEK.	¢19
hukas	NGS - RIK L	505
×1HA5	HOTSGAKK INGHIMPI LPLSINA AVECGOVGYS LYMDOQNOWS	533
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hunas		546
£18A5	TPEKOREHY- WHILTGOVGY VHYHVIHAVH YHVHVKR CCR-KRSQTV	527
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FIG. 2

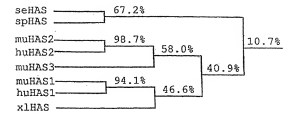


FIG. 3

SIZE DISTRIBUTION OF HYALURONAN PRODUCED BY DIFFERENT ENGINEERED STREPTOCOCCAL HAS ENZYMES

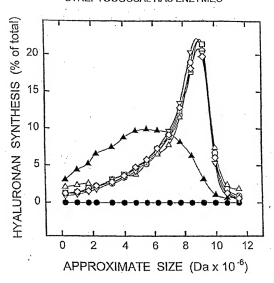


FIG. 4

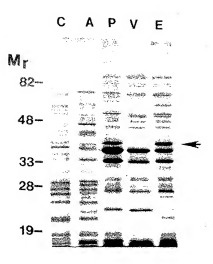


Fig. 5

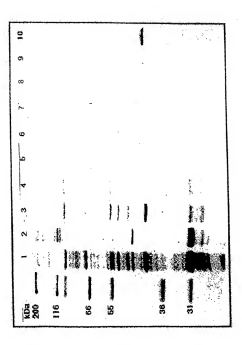


Fig. 6

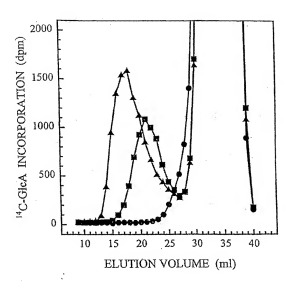
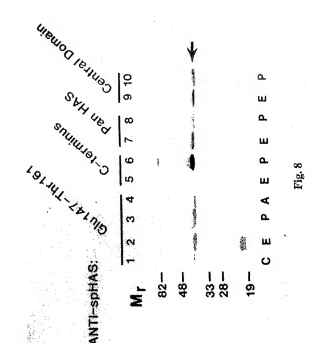


FIG. 7



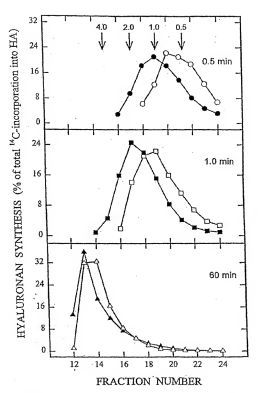


FIG. 9

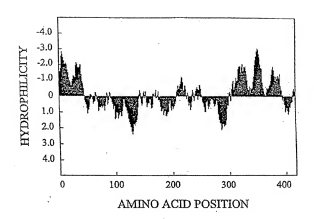


FIG. 10

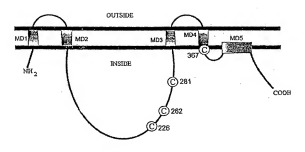
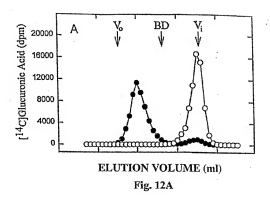
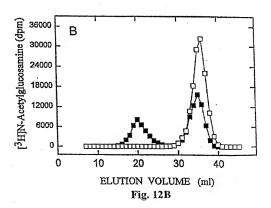
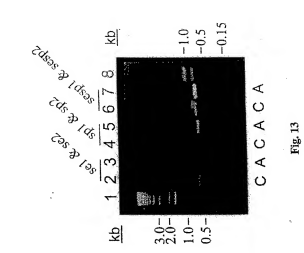


FIG. 11







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+ sesp2	1254

FIG. 14